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$%\STN;HighlightOn= ***;HighlightOff=***
Connecting via Winsock to STN
welcome to STN International! Enter x:x
LOGINID:SSSPTA1635KXH
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TERMINAL (ENTER 1, 2, 3, OR ?):2
* * * * * * * * * *
                      Welcome to STN International
                  Web Page URLs for STN Seminar Schedule - N. America
 NEWS
       1
                   'Ask CAS" for self-help around the clock
 NEWS
                   Source of Registration (SR) information in REGISTRY updated
          JAN 27
 NEWS
                   and searchable
                   A new search aid, the Company Name Thesaurus, available in
 NEWS
       4
          JAN 27
                   CA/CAplus
          FEB 05
                  German (DE) application and patent publication number format
 NEWS
       5
                   changes
          MAR 03
                  MEDLINE and LMEDLINE reloaded
 NEWS
       6
          MAR 03
                  MEDLINE file segment of TOXCENTER reloaded
 NEWS
 NEWS
          MAR 03
                   FRANCEPAT now available on STN
          MAR 29
                  Pharmaceutical Substances (PS) now available on STN
 NEWS
 NEWS 10
          MAR 29
                  WPIFV now available on STN
          MAR 29
                  New monthly current-awareness alert (SDI) frequency in RAPRA
 NEWS
      11
                  PROMT: New display field available IFIPAT/IFIUDB/IFICDB: New super search and display field
          APR 26
 NEWS
      12
 NEWS 13
          APR 26
                   available
          APR 26
                   LITALERT now available on STN
 NEWS 14
          APR 27
                   NLDB: New search and display fields available
 NEWS 15
 NEWS 16
          May 10
                   PROUSDDR now available on STN
 NEWS 17
          May 19
                   PROUSDDR: One FREE connect hour, per account, in both May
                   and June 2004
                  EXTEND option available in structure searching Polymer links for the POLYLINK command completed in REGISTRY
 NEWS 18
          May 12
 NEWS 19
          May 12
          May 17
 NEWS .20
                   FRFULL now available on STN
                   STN User Update to be held June 7 and June 8 at the SLA 2004
 NEWS 21
          May 27
                   Conference
                  New UPM (Update Code Maximum) field for more efficient patent
 NEWS 22
          May 27
                   SDIs in CAplus
                   CAplus super roles and document types searchable in REGISTRY
 NEWS 23
          May 27
 NEWS 24
          May 27
                  Explore APOLLIT with free connect time in June 2004
               MARCH 31 CURRENT WINDOWS VERSION IS V7.00A, CURRENT
 NEWS EXPRESS
               MACINTOSH VERSION IS V6.0c(ENG) AND V6.0Jc(JP)
               AND CURRENT DISCOVER FILE IS DATED 26 APRIL 2004
 NEWS HOURS
               STN Operating Hours Plus Help Desk Availability
               General Internet Information
 NEWS INTER
 NEWS LOGIN
               Welcome Banner and News Items
               Direct Dial and Telecommunication Network Access to STN
 NEWS PHONE
               CAS world wide web Site (general information)
 NEWS WWW
Enter NEWS followed by the item number or name to see news on that
specific topic.
  All use of STN is subject to the provisions of the STN Customer
  agreement. Please note that this agreement limits use to scientific
  research. Use for software development or design or implementation of commercial gateways or other similar uses is prohibited and may
  result in loss of user privileges and other penalties.
  FILE 'HOME' ENTERED AT 12:55:07 ON 08 JUN 2004
=> s cytomegalovirus or (herpes simplex) or (varicellar)
THIS COMMAND NOT AVAILABLE IN THE CURRENT FILE
Some commands only work in certain files. For example, the EXPAND
command can only be used to look at the index in a file which has an
        Enter "HELP COMMANDS" at an arrow prompt (=>) for a list of
index.
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=> b medline caplus lifesci embase uspatfull biosis
COST IN U.S. DOLLARS
                                                      SINCE FILE
                                                                       TOTAL
                                                           ENTRY
                                                                      SESSION
FULL ESTIMATED COST
                                                                         0.42
                                                             0.42
FILE 'MEDLINE' ENTERED AT 12:56:01 ON 08 JUN 2004
FILE 'CAPLUS' ENTERED AT 12:56:01 ON 08 JUN 2004
USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.
PLEASE SEE "HELP USAGETERMS" FOR DETAILS.
COPYRIGHT (C) 2004 AMERICAN CHEMICAL SOCIETY (ACS)
FILE 'LIFESCI' ENTERED AT 12:56:01 ON 08 JUN 2004
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FILE 'EMBASE' ENTERED AT 12:56:01 ON 08 JUN 2004 COPYRIGHT (C) 2004 Elsevier Inc. All rights reserved.
FILE 'USPATFULL' ENTERED AT 12:56:01 ON 08 JUN 2004
CA INDEXING COPYRIGHT (C) 2004 AMERICAN CHEMICAL SOCIETY (ACS)
FILE 'BIOSIS' ENTERED AT 12:56:01 ON 08 JUN 2004
COPYRIGHT (C) 2004 BIOLOGICAL ABSTRACTS INC. (R)
=> s cytomegalovirus or (herpes simplex) or (varicellar)
         235427 CYTOMEGALOVIRUS OR (HERPES SIMPLEX) OR (VARICELLAR)
L1
=> s l1 and (calpain or e64D) or (z()leu()leu()h)
            340 L1 AND (CALPAIN OR E64D) OR (Z(W) LEU(W) LEU(W) H)
=> dup rem 12
PROCESSING COMPLETED FOR L2
             323 DUP REM L2 (17 DUPLICATES REMOVED)
\Rightarrow s 11 and ((calpain or e64D) or (z()leu()leu()h))
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PROCESSING COMPLETED FOR L4
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   3 FILES SEARCHED...
             26 L5 AND PY<2000
=> d 16 ibib abs tot
     ANSWER 1 OF 26 CAPLUS COPYRIGHT 2004 ACS on STN
                            1999:723196 CAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                            131:333006
                           Production of recombinant replication-deficient viral
TITLE:
                           vectors encoding exogenous transgenes via
                           microcarrier-based process
INVENTOR(S):
                           Giroux, Daniel D.; Goudreau, Ann M.; Ramachandra,
                           Muralidhara; Shabram, Paul W.
                           Canji, Inc., USA
PCT Int. Appl., 32 pp.
PATENT ASSIGNEE(S):
SOURCE:
                           CODEN: PIXXD2
DOCUMENT TYPE:
                            Patent
LANGUAGE:
                            English
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                                               APPLICATION NO.
     PATENT NO.
                        KIND DATE
                                                                  DATE
                                               wo 1999-US9813
                                                                 19990504 <--
     wo 9957297
                         Α1
                               19991111
         W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CZ, DE,
              DK, EE, ES, FI, GB, GD, GE, HR, HU, ID, IL, IN, IS, JP, KG, KR,
         RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                                               ús 1998-73076
     us 5994134
                               19991130
                                                                  19980504 <--
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AU 9938823
                                    19991123
                                                       AU 1999-38823
                             Α1
                                                                              19990504 <--
                                    20010228
                                                       EP 1999-921681
      EP 1078095
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                                                                             19990504
                AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, PT, IE,
                 LT, LV, FI, RO
      JP 2002513583
                             T2
                                    20020514
                                                       JP 2000-547250
                                                                             19990504
                                                   US 1998-73076 A 19980504
WO 1999-US9813 W 19990504
PRIORITY APPLN. INFO.:
      The present invention is directed to a method of producing recombinant viral vectors at high titers incorporating a variety of important advancements over the art. The method of the present invention
      incorporates multiple features which provide enhanced prodn. of viruses,
      particularly those viruses encoding exogenous transgenes. The
      specifically illustrated method describes a method for the high titer
      serum-free media prodn. of recombinant replication defective adenoviruses
      contg. an exogenous transgene. The invention provides methods of prepg. microcarriers, methods for seeding bioreactors at high cell d., increasing the infectivity of the producer cells to the virus, methods to increase product yield through synchronization of the cell cycle of the producer
      cells, and methods to minimize the deleterious effects of exogenous
      transgenes. The invention further provides producer cells prepd. by the
      process of the invention. The invention further provides viruses produced
      by the process.
REFERENCE COUNT:
                                5
                                        THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS
                                        RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
      ANSWER 2 OF 26 CAPLUS COPYRIGHT 2004 ACS on STN
                                1999:718875 CAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                                131:348774
TITLE:
                                Tandem fluorescent protein constructs and their
                                preparation for enzyme assays
INVENTOR(S):
                                Tsien, Roger Y.; Heim, Roger; Cubitt, Andrew
PATENT ASSIGNEE(S):
                                The Regents of the University of California, USA;
                                Aurora Biosciences Corporation
SOURCE:
                                U.S., 33 pp., Cont.-in-part of U.S. Ser. No. 594,575.
                                CODEN: USXXAM
DOCUMENT TYPE:
                                Patent
LANGUAGE:
                                English
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
      PATENT NO.
                            KIND
                                    DATE
                                                       APPLICATION NO.
                                                                             DATE
      US 5981200
                                    19991109
                                                       US 1997-792553
                                                                             19970131 <--
      PT 877805
                                    20021031
                                                       PT 1997-905667
                             Т
                                                                             19970131
                                    20021216
                                                       ES 1997-905667
      ES 2177939
                             Т3
                                                                              19970131
      us 2003186229
                                    20031002
                             Α1
                                                       US 2001-865291
                                                                              20010524
      US 2002164674
                                    20021107
                                                       US 2002-57505
                             Α1
                                                                              20020125
                                                   us 1996-594575
us 1997-792553
                                                                        A2 19960131
A1 19970131
PRIORITY APPLN. INFO.:
                                                   us 1999-396003
                                                                         в2 19990913
      This invention provides tandem fluorescent protein construct including a
      donor fluorescent protein moiety, an acceptor fluorescent protein moiety and a linker moiety that couples the donor and acceptor moieties. The
      donor and acceptor moieties exhibit fluorescence resonance energy transfer
      which is eliminated upon cleavage. The constructs are useful in enzymic assays. Mutant green fluorescent proteins (GFPs) were created by mutagenesis of the Aequorea victoria GFP. Polyhistidine tagged tandem green and blue fluorescent proteins were recombinantly constructed having an inserted peptide sequence including cleavage recognition sites for many
      proteases. Cleavage expts. were done with trypsin, enterokinase and
         ***calpain***
REFERENCE COUNT:
                                22
                                        THERE ARE 22 CITED REFERENCES AVAILABLE FOR THIS
                                        RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
      ANSWER 3 OF 26 CAPLUS COPYRIGHT 2004 ACS on STN
                                1998:728567 CAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                                130:10614
JITLE:
                                Ricin precursors cleavable by disease-specific
                                proteinases for treatment of cancer, viral or
                                parasitic infections
Borgford, Thor
    NTOR(S):
       ASSIGNEE(S):
                                De Novo Enzyme Corp., Can.
                                PCT Int. Appl., 352 pp.
                                CODEN: PIXXD2
             YPE:
                                Patent
                                English
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AB

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KIND DATE
      PATENT NO.
                                                  APPLICATION NO.
                                                                       DATE
      wo 9849311
                                 19981105
                                                                       19980430 <---
                          Α2
                                                  WO 1998-CA394
      wo 9849311
                          Α3
                                 19990211
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          W:
               KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX,
               NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT,
               UA, UG, US, UZ, VN, YU, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
          RW: GH, GM, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG
      AU 9870237
      EP 977862
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               AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
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                                                                       19980430
                                                  us 1999-403752
      us 6593132
                                                                       19991029
                                 20030715
                           В1
      us 2004009551
                          Α1
                                 20040115
                                                  US 2003-394511
                                                                       20030324
                                               US 1997-45148P
PRIORITY APPLN. INFO.:
                                                                  Р
                                                                       19970430
                                               US 1997-63715P
                                                                       19971029
                                                                   Ρ
                                               WO 1998-CA394
                                                                   W 19980430
                                               US 1999-403752
                                                                 A3 19991029
      Ricin precursors with the ricin A and B chains linked by a protease-labile
AB
      linker peptide are described for use in the treatment of disease. The
      linker peptide contains a cleavage site for a disease specific protease
      such as a cancer, fungal, viral or parasitic protease. The ricin A chain
     may be replaced by comparable cytotoxic proteins such as the abrin A chain. The protein is delivered to the target tissue using viral vectors carrying an expression cassette for the ricin fusion protein gene.
      Construction of a series of variants of preproricin cleavable by a no. of
      different proteinases is described. Cleavage and activation of these
      variants with the expected patterns of cleavage of rRNA is demonstrated.
     ANSWER 4 OF 26 USPATFULL on STN
L6
ACCESSION NUMBER:
                            2003:296940 USPATFULL
TITLE:
                            Lactacystin analogs
                            Schreiber, Stuart L., Boston, MA, United States
Standaert, Robert F., Bryan, TX, United States
INVENTOR(S):
                            Fenteany, Gabriel, Cambridge, MA, United States
Jamison, Timothy F., Cambridge, MA, United States
PATENT ASSIGNEE(S):
                            Millennium Pharmaceuticals, Inc., Cambridge, MA, United
                            States (U.S. corporation)
                                  NUMBER
                                                 KIND
                                                          DATE
PATENT INFORMATION:
                            us 6645999
                                                        20031111
                                                  в1
                            wo 9632105
                                                         19961017
                            US 1997-945092
APPLICATION INFO.:
                                                         19970126
                                                                    (8)
                            wo 1996-US5072
                                                         19960412
                                                         19980126 PCT 371 date
RELATED APPLN. INFO.:
                            Continuation-in-part of Ser. No. US 1995-421583, filed
                            on 12 Apr 1995
                            Utility
DOCUMENT TYPE:
FILE SEGMENT:
                            GRANTED
PRIMARY EXAMINER:
                            Travers, Russell
LEGAL REPRESENTATIVE:
                            Hale and Dorr LLP
NUMBER OF CLAIMS:
EXEMPLARY CLAIM:
NUMBER OF DRAWINGS:
                            0 Drawing Figure(s); 0 Drawing Page(s)
LINE COUNT:
                            2868
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
AΒ
        Compounds related to lactacystin and lactacystin Beta-lactone
        pharmaceutical compositions containing the compounds, and methods of
        use.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
```

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L6 ANSWER 5 OF 26
ACCESSION NUMBER: 2003:190684 USPATFULL
TITLE: Ricin-like toxin variants for treatment of cancer, viral or parasitic infections
INVENTOR(S): Borgford, Thor, Burnaby, CANADA
PATENT ASSIGNEE(S): Twinstrand Therapeutics Inc., Vancouver, CANADA
```

NUMBER KIND DATE US 6593132 PATENT INFORMATION: 20030715 в1 WO 9849311 US 1999-403752 19981105 <--APPLICATION INFO.: 19991029 (9)WO 1998-CA394 19980430 OCUMENT TYPE: Utility FILE SEGMENT: **GRANTED** PRIMARY EXAMINER: Carlson, Karen Cochrane \_EGAL REPRESENTATIVE: Bereskin & Parr, Gravelle, Micheline NUMBER OF CLAIMS: EXEMPLARY CLAIM: NUMBER OF DRAWINGS: 254 Drawing Figure(s); 254 Drawing Page(s) INE COUNT: 5176
CAS INDEXING IS AVAILABLE FOR THIS PATENT. The present invention provides a protein having an A chain of a ۱В ricin-like toxin, a B chain of a ricin-like toxin and a heterologous linker amino acid sequence, linking the A and B chains. The linker sequence contains a cleavage recognition site for a disease specific protease such as a cancer, fungal, viral or parasitic protease. The invention also relates to a nucleic acid molecule encoding the protein and to expression vectors incorporating the nucleic acid molecule. Also provided is a method of inhibiting or destroying mammalian cancer cells, cells infected with a virus, a fungus, or parasite, or parasites utilizing the nucleic acid molecules and proteins of the invention and pharmaceutical compositions for treating human cancer, viral infection pharmaceutical compositions for treating human cancer, viral infection, fungal infection, or parasitic infection. CAS INDEXING IS AVAILABLE FOR THIS PATENT. ANSWER 6 OF 26 USPATFULL ON STN 2002:115819 USPATFULL ACCESSION NUMBER: TITLE: Fibrinogen-coated particles for therapeutic use INVENTOR(S): Yen, Richard C. K., Yorba Linda, CA, United States Hemosphere, Inc., Anaheim, CA, United States (U.S. PATENT ASSIGNEE(S): corporation) NUMBER KIND DATE PATENT INFORMATION: US 6391343 20020521 в1 wo 9639128 19961212 <--US 1998-952765 APPLICATION INFO.: 19980410 (8) wo 1996-us9458 19960604 19980410 PCT 371 date Continuation-in-part of Ser. No. US 1995-554919, filed on 9 Nov 1995, now abandoned Continuation-in-part of Ser. No. US 1995-471650, filed on 6 Jun 1995, now patented, Pat. No. US 5725804 Continuation-in-part of Ser. No. US 1994-212546, filed on 14 Mar 1994, now RELATED APPLN. INFO.: patented, Pat. No. US 5616311 Continuation-in-part of Ser. No. US 1993-69831, filed on 1 Jun 1993, now abandoned Continuation-in-part of Ser. No. US 1992-959560, filed on 13 Oct 1992, now patented, Pat. No. US 5308620 Continuation-in-part of Ser. No. US 1991-641720, filed on 15 Jan 1991, now abandoned Utility OCUMENT TYPE: ILE SEGMENT: **GRANTED** RIMARY EXAMINER: Lovering, Richard D. EGAL REPRESENTATIVE: Townsend and Townsend and Crew LLP NUMBER OF CLAIMS: 11 XEMPLARY CLAIM: UMBER OF DRAWINGS: 0 Drawing Figure(s); 0 Drawing Page(s) INE COUNT: 2407 AS INDEXING IS AVAILABLE FOR THIS PATENT.

B The invention provides a particle comprising fibrinogen bound on the

by it self of formation of a clot upon activation by thrombin.

AS INDEXING IS AVAILABLE FOR THIS PATENT.

6 ANSWER 7 OF 26 USPATFULL ON STN

CCESSION NUMBER: 2001:82299 USPATFULL

ITLE: Method and product for cleaning and/or whitening of

surface of an albumin matrix, wherein said particle is capable of

coaggregation with platelet, and of aggregation in a solution containing soluble fibrinogen at a concentration of soluble fibrinogen not capable

Rinne, Ari, Pajutie 3 B, FIN-2G900 Turku, Finland VENTOR(S):

NUMBER KIND DATE us 6241973 20010605 TENT INFORMATION: в1 wo 9829088 19980709 us 1999-331777 PLICATION INFO.: 19990624 (9) wo 1998-FI1 19980102

19990624 PCT 371 date 19990624 PCT 102(e) date

<--

NUMBER DATE ·----

IORITY INFORMATION: FI 1997-12 19970103

Utility CUMENT TYPE: LE SEGMENT: Granted IMARY EXAMINER:

Rose, Shep K. Lydon, James C.

MBER OF CLAIMS: EMPLARY CLAIM: NE COUNT:

GAL REPRESENTATIVE:

VENTOR(S):

TENT ASSIGNEE(S):

LATED APPLN. INFO.:

S INDEXING IS AVAILABLE FOR THIS PATENT.

A method and a product for cleaning and/or whitening of teeth. Natural human cysteine proteinases are employed for cleaning and whitening purposes and this activity can be blocked by natural cysteine protease inhibitors, which are released secondarily from the product at a later stage. The use of natural cysteine proteinases and their inhibitors provides the advantage that they are man's own proteins, and therefore the risk of allegorization is minimized. In addition, their enzyme kinetics are will known.

## S INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 8 OF 26 USPATFULL on STN

CESSION NUMBER: 2000:41033 USPATFULL

Synthetic catalytic free radical scavengers useful as TLE:

antioxidants for prevention and therapy of disease Malfroy-Camine, Bernard, Arlington, MA, United States Doctrow, Susan Robin, Roslindale, MA, United States

Eukarion, Inc., Bedford, MA, United States (U.S.

corporation)

NUMBER KIND DATE US 6046188 20000404 TENT INFORMATION: wo 9640148 19961219 <--US 1998-973577 19980311 (8) PLICATION INFO.: wo 1996-us10037 19960606 19980311 PCT 371 date

19980311 PCT 102(e) date Continuation-in-part of Ser. No. US 1995-485489, filed

on 7 Jun 1995, now patented, Pat. No. US 5696109 Utility CUMENT TYPE: Granted LE SEGMENT:

Reamer, James H. Townsend & Townsend & Crew LLP IMARY EXAMINER:

GAL REPRESENTATIVE:

MBER OF CLAIMS: EMPLARY CLAIM:

MBER OF DRAWINGS: NE COUNT:

28 Drawing Figure(s); 16 Drawing Page(s) 3405

S INDEXING IS AVAILABLE FOR THIS PATENT.

The invention provides antioxidant salen-metal complexes, compositions of such antioxidant salen-metal complexes having superoxide activity, catalase activity, and/or peroxidase activity, compositions of salen-metal complexes in a form suitable for pharmaceutical administration to treat or prevent a disease associated with cell or tissue damage produced by free radicals such as superoxide, and cosmetic and free radical quenching formulations of salen metal compounds.

### S INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 9 OF 26 USPATFULL on STN

1999:166974 USPATFULL CESSION NUMBER:

TLE: Cysteine protease inhibitors VENTOR(S):

Spruce, Lyle W., Chula Vista, CA, United States Gyorkos, Albert C., Westminster, CO, United States

Goodfellow, Val S., Tucson, AZ, United States Leimer, Axel H., Westborough, MA, United States Young, John M., Redwood City, CA, United States Gerrity, James Ivan, Albany, OR, United States Cortech Inc., Bedminster, NJ, United States (U.S.

PATENT ASSIGNEE(S): corporation)

> NUMBER KIND DATE

PATENT INFORMATION: APPLICATION INFO.:

us 6004933 19991221 <-us 1998-65258 19980423 (9)

> NUMBER DATE

PRIORITY INFORMATION:

US 1997-44819P 19970425 (60)

DOCUMENT TYPE: FILE SEGMENT:

Utility

Granted

PRIMARY EXAMINER: ASSISTANT EXAMINER: Richter, Johann Solola, Taofiq A. Dechert Price & Rhoads

LEGAL REPRESENTATIVE:

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

146

NUMBER OF DRAWINGS:

4 Drawing Figure(s); 3 Drawing Page(s)

LINE COUNT:

2591

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention relates to cysteine protease inhibitors of the general formula (I): ##STR1## wherein Z is a cysteine protease binding moiety; X and Y are S, O or optionally substituted N; and R.sub.1 is optionally substituted alkyl or aryl.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 10 OF 26 USPATFULL on STN L6

1999:155518 USPATFULL

ACCESSION NUMBER: TITLE:

Viral production process

INVENTOR(S):

Giroux, Daniel D., La Jolla, CA, United States Goudreau, Ann M., San Diego, CA, United States Ramachandra, Muralidhara, San Diego, CA, United States

PATENT ASSIGNEE(S):

Shabram, Paul W., Olivenhain, CA, United States Canji, Inc., San Diego, CA, United States (U.S.

corporation)

NUMBER KIND DATE

PATENT INFORMATION:

APPLICATION INFO.:

us 5994134 us 1998-73076 19991130 19980504 (9)

DOCUMENT TYPE: FILE SEGMENT:

Utility Granted

PRIMARY EXAMINER:

Stucker, Jeffrey Murphy, Richard B.

LEGAL REPRESENTATIVE:

NUMBER OF CLAIMS:

17

EXEMPLARY CLAIM: NUMBER OF DRAWINGS:

3 Drawing Figure(s); 3 Drawing Page(s)

LINE COUNT:

1005

CAS INDEXING IS AVAILABLE FOR THIS PATENT. AΒ

The present invention is directed to a method of producing recombinant viral vectors at high titers incorporating a variety of important advancements over the art. The method of the present invention incorporates multiple features which provide enhanced production of viruses, particularly those viruses encoding exogenous transgenes. The specifically illustrated method describes a method for the high titer serum-free media production of recombinant replication defective adenoviruses containing an exogenous transgene. The invention provides methods of preparing microcarriers, methods for seeding bioreactors at high cell density, increasing the infectivity of the producer cells to the virus, methods to increase product yield through synchronization of the cell cycle of the producer cells, and methods to minimize the deleterious effects of exogenous transgenes. The invention further provides producer cells prepared by the process of the invention. The invention further provides viruses produced by the process.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 11 OF 26 USPATFULL on STN L6

ACCESSION NUMBER: 1999:137014 USPATFULL TITLE:

Vesicle transport related protein

Corley, Neil C., Mountain View, CA, United States

Shah, Purvi, Sunnyvale, CA, United States Incyte Pharmaceuticals, Inc., Palo Alto, CA, United

States (U.S. corporation)

NUMBER KIND DATE PATENT INFORMATION: us 5976865 19991102

APPLICATION INFO.: Utility DOCUMENT TYPE:

us 1997-984172

19971203 (8)

<--

FILE SEGMENT:

Granted

PRIMARY EXAMINER:

Johnson, Nancy A. Incyte Pharmaceuticals, Inc.

\_EGAL REPRESENTATIVE: NUMBER OF CLAIMS: EXEMPLARY CLAIM:

10

7 Drawing Figure(s); 7 Drawing Page(s)

NUMBER OF DRAWINGS: LINE COUNT:

PATENT ASSIGNEE(S):

2242

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

٩В

The invention provides a human vesicle transport related protein (VTRP) and polynucleotides which identify and encode VTRP. The invention also provides expression vectors, host cells, antibodies, agonists, and antagonists. The invention also provides methods for treating or preventing disorders associated with expression of VTRP.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 12 OF 26 USPATFULL ON STN SSION NUMBER: 1999:102423 USPATFULL -6

ACCESSION NUMBER:

TITLE:

PATENT ASSIGNEE(S):

Method for making non-crosslinked protein particles for

therapeutic and diagnostic use

INVENTOR(S):

Yen, Richard C. K., Glendora, CA, United States Hemosphere, Inc., Irvine, CA, United States (U.S.

corporation)

KIND DATE NUMBER

PATENT INFORMATION: APPLICATION INFO.:

us 5945033 19990831 US 1996-747137 19961112

RELATED APPLN. INFO.:

Continuation of Ser. No. US 1994-212546, filed on 14 Mar 1994, now patented, Pat. No. US 5616311 which is a continuation-in-part of Ser. No. US 1993-69831, filed

on 1 Jun 1993, now abandoned And Ser. No. US 1993-69831, filed on 1 Jun 1993, now abandoned And Ser. No. US 1992-959560, filed on 13 Oct 1992, now patented, Pat. No. US 5308620 which is a continuation-in-part of Ser. No. US 1991-641720, filed on 15 Jan 1991, now abandoned Utility

DOCUMENT TYPE: FILE SEGMENT:

Granted Dees, Jose' G.

PRIMARY EXAMINER: ASSISTANT EXAMINER: LEGAL REPRESENTATIVE:

Hartley, Michael G. Townsend and Townsend and Crew LLP

NUMBER OF CLAIMS:

12

EXEMPLARY CLAIM: \_INE COUNT:

3655

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB Albumin particles in the nanometer and micrometer size range in an aqueous suspension are rendered stable against resolubilization without the aid of a crosslinking agent and without denaturation, by the incorporation of hemoglobin in the particle composition. Particles which are primarily hemoglobin in the nanometer and micrometer size range in an aqueous suspension are rendered stable against aggregation by the incorporation of either albumin, surface active agents or gelatin.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 13 OF 26 USPATFULL ON STN

ACCESSION NUMBER:

1999:81758 USPATFULL

TITLE:

Non-activated receptor complex proteins and uses

thereof

INVENTOR(S):

Davis, Roger J., Princeton, MA, United States

Galcheva-Gargova, Zoya, Worcester, MA, United States University of Massachusetts, Boston, MA, United States

(U.S. corporation)

NUMBER KIND DATE \_\_\_\_\_

PATENT INFORMATION:

PATENT ASSIGNEE(S):

us 5925566

19990720

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NUMBER DATE
PRIORITY INFORMATION:
                             US 1996-19219P 19960606 (60)
DOCUMENT TYPE:
                             Utility
FILE SEGMENT:
                             Granted
PRIMARY EXAMINER:
                             Campell, Bruce R.
                             Nguyen, Dave Trong
ASSISTANT EXAMINER:
                             Fish & Richardson, P.C.
LEGAL REPRESENTATIVE:
                             23
NUMBER OF CLAIMS:
EXEMPLARY CLAIM:
NUMBER OF DRAWINGS:
                             22 Drawing Figure(s); 18 Drawing Page(s)
                             2438
LINE COUNT:
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
        The invention features a substantially pure ZPR1 polypeptide. For example, a ZPR1 polypeptide that specifically binds to a non-activated
AΒ
        membrane-bound receptor (e.g., EGF or PDGF receptors) and specifically
        binds small nucleolar RNAs (e.g., U3). ZPR1 polypeptides can be isolated from any eukaryote, including mammals (e.g. rodents and humans) and
        fungi (e.g., S. cerevisiae and S. pombe).
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
     ANSWER 14 OF 26 USPATFULL on STN
ACCESSION NUMBER:
                             1999:36949 USPATFULL
TITLE:
                             Engineering oral tissues
INVENTOR(S):
                             Mooney, David J., Ann Arbor, MI, United States
                            Rutherford, Robert B., Ann Arbor, MI, United States
The Regents of the University of Michigan, Ann Arbor,
MI, United States (U.S. corporation)
PATENT ASSIGNEE(S):
                                  NUMBER
                                                 KIND
                                                          DATE
                            US 5885829
US 1997-864494
PATENT INFORMATION:
                                                          19990323
                                                                                   <--
APPLICATION INFO.:
                                                          19970528 (8)
                                     NUMBER DATE
PRIORITY INFORMATION:
                             US 1996-18450P 19960528 (60)
DOCUMENT TYPE:
                             Utility
FILE SEGMENT:
                             Granted
PRIMARY EXAMINER:
                             Degen, Nancy
LEGAL REPRESENTATIVE:
                             Arnold, White & Durkee
NUMBER OF CLAIMS:
                             109
EXEMPLARY CLAIM:
NUMBER OF DRAWINGS:
                             17 Drawing Figure(s); 11 Drawing Page(s)
LINE COUNT:
                             8001
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       Disclosed are methods for regenerating dental and oral tissues from
٩B
        viable cells using ex vivo culture on a structural matrix. The
        regenerated oral tissues and tissue-matrix preparations thus provided
       have both clinical applications in dentistry and oral medicine and are also useful in in vitro toxicity and biocompatibility testing.
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
L6
    ANSWER 15 OF 26 USPATFULL ON STN
ACCESSION NUMBER:
                             1998:138941 USPATFULL
TTTLF:
                             Synthetic catalytic free radical scavengers useful as
                            antioxidants for prevention and therapy of disease
Malfroy-Camine, Bernard, Arlington, MA, United States
Doctrow, Susan Robin, Roslindale, MA, United States
INVENTOR(S):
PATENT ASSIGNEE(S):
                             Eukarion, Inc., Bedford, MA, United States (U.S.
                             corporation)
                                 NUMBER KIND DATE
                            us 5834509
us 1995-479697
PATENT INFORMATION:
                                                          19981110
                            US 1995-479697 19950607 (8)
Continuation-in-part of Ser. No. US 1995-380731, filed on 26 Jan 1995 which is a continuation-in-part of Ser.
APPLICATION INFO.:
RELATED APPLN. INFO.:
                            No. US 1992-987474, filed on 7 Dec 1992, now patented,
                             Pat. No. US 5403834
```

NUMBER DATE

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

Jarvis, William R. A. PRIMARY EXAMINER:

LEGAL REPRESENTATIVE: Townsend and Townsend and Crew LLP

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 28 Drawing Figure(s); 19 Drawing Page(s)

LINE COUNT: 3384

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

This invention provides salen-manganese complexes and pharmaceutically AB acceptable compositions thereof useful as antioxidants and free radical

scavengers.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 16 OF 26 USPATFULL on STN

ACCESSION NUMBER: 1998:134999 USPATFULL

Methods for the treatment of bone resorption disorders, TITLE:

including osteoporosis

INVENTOR(S):

Gelb, Bruce D., Dobbs Ferry, NY, United States Chapman, Harold, Newton, MA, United States Desnick, Robert J., New York, NY, United States Mount Sinai School of Medicine of the City of New York,

PATENT ASSIGNEE(S): New York, NY, United States (U.S. corporation)

Brigham and Women's Hospital, Boston, MA, United States

<--

<--

(U.S. corporation)

NUMBER KIND DATE

PATENT INFORMATION: APPLICATION INFO.:

us 5830850 19981103 us 1996-704473 19960828 (8)

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

PRIMARY EXAMINER: Marschel, Ardin H. LEGAL REPRESENTATIVE: Pennie & Edmonds LLP

NUMBER OF CLAIMS: **EXEMPLARY CLAIM:** 

NUMBER OF DRAWINGS: 7 Drawing Figure(s); 6 Drawing Page(s)

LINE COUNT: 2434

CAS INDEXING IS AVAILABLE FOR THIS PATENT. AΒ

The present invention relates to methods and compositions for the amelioration of symptoms caused by bone resorption disorders, including but not limited to osteoporosis, arthritides and periodontal disease, and damage caused by macrophage-mediated inflammatory processes. In one embodiment, the methods and compositions of the invention include methods and compositions for the specific inhibition of cathepsin  $\kappa$ activity. In an additional embodiment, the methods and compositions of the invention include methods and compositions for the specific inhibition of cathepsin K activity coupled with specific inhibition of at least a second activity involved in the bone resorption and/or macrophage-mediated inflammatory processes. In a particular embodiment, the methods and compositions of the invention include methods and compositions for the specific inhibition of cathepsin K and cathepsin S activity.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 17 OF 26 USPATFULL on STN

ACCESSION NUMBER: 1998:131743 USPATFULL

TITLE: Synthetic catalytic free radical scavengers useful as

antioxidants for prevention and therapy of disease Malfroy-Camine, Bernard, Arlington, MA, United States

Doctrow, Susan Robin, Roslindale, MA, United States

PATENT ASSIGNEE(S): Eukarion, Inc., Bedford, MA, United States (U.S.

corporation)

KIND NUMBER DATE

US 5827880 US 1995-380731 PATENT INFORMATION: 19981027 APPLICATION INFO.:

19950126 (8)

Continuation-in-part of Ser. No. US 1992-987474, filed RELATED APPLN. INFO.:

on 7 Dec 1992, now patented, Pat. No. US 5403834

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

INVENTOR(S):

PRIMARY EXAMINER: Nazario-Gonzalez, Porfirio

LEGAL REPRESENTATIVE: Townsend and Townsend and Crew LLP EXEMPLARY CLAIM: 1,12

28 Drawing Figure(s); 19 Drawing Page(s) NUMBER OF DRAWINGS:

LINE COUNT: 3241

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The invention provides antioxidant salen-metal complexes, compositions of such antioxidant salen-metal complexes having superoxide activity,

catalase activity, and/or peroxidase activity, compositions of salen-metal complexes in a form suitable for pharmaceutical

administration to treat or prevent a disease associated with cell or tissue damage produced by free radicals such as superoxide, and cosmetic

and free radical quenching formulations of salen metal compounds.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 18 OF 26 USPATFULL ON STN

ACCESSION NUMBER: 1998:58182 USPATFULL Lactacystin analogs TITLE:

Fenteany, Gabriel, Cambridge, MA, United States Jamison, Timothy F., Cambridge, MA, United States INVENTOR(S):

Schreiber, Stuart L., Boston, MA, United States

Standaert, Robert F., Arlington, MA, United States President and Fellows of Harvard College, Cambridge, PATENT ASSIGNEE(S):

MA, United States (U.S. corporation)

NUMBER KIND DATE

us 5756764 PATENT INFORMATION: 19980526

us 1995-466468 19950606 APPLICATION INFO.: (8)

Division of Ser. No. US 1995-421583, filed on 12 Apr RELATED APPLN. INFO.:

1995

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

PRIMARY EXAMINER: Richter, Johann ASSISTANT EXAMINER: Stockton, Laura L. LEGAL REPRESENTATIVE: Fish & Richardson P.C.

NUMBER OF CLAIMS: 16 EXEMPLARY CLAIM: LINE COUNT: 2392

AR

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Described herein are compounds related to lactacystin and lactacystin .beta.-lactone, pharmaceutical compositions containing the compounds,

and methods of use.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 19 OF 26 USPATFULL ON STN SSION NUMBER: 1998:57716 USPATFULL ACCESSION NUMBER:

Aptamers specific for biomolecules and methods of TITLE:

making

INVENTOR(S): Griffin, Linda, Atherton, CA, United States

Albrecht, Glenn, Redwood City, CA, United States

Latham, John, Palo Alto, CA, United States

Leung, Lawrence, Hillsborough, CA, United States Vermaas, Eric, Oakland, CA, United States Toole, John J., Burlingame, CA, United States Gilead Sciences, Inc., Foster City, CA, United States

PATENT ASSIGNEE(S):

(U.S. corporation)

NUMBER KIND DATE

US 5756291 US 1995-484192 PATENT INFORMATION: 19980526 <--

19950607 APPLICATION INFO.: Continuation of Ser. No. US 1992-934387, filed on 21 RELATED APPLN. INFO.:

Aug 1992, now abandoned

Utility DOCUMENT TYPE: FILE SEGMENT: Granted

PRIMARY EXAMINER: Zitomer, Stephanie W.

LEGAL REPRESENTATIVE: Bosse, Mark L.

NUMBER OF CLAIMS: 12 **EXEMPLARY CLAIM:** 

NUMBER OF DRAWINGS: 6 Drawing Figure(s); 6 Drawing Page(s)

LINE COUNT: 8242

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

A method for identifying oligomer sequences, optionally comprising modified base, which specifically bind target molecules such as serum proteins, kinins, eicosanoids and extracellular proteins is described.

PDGF, FGF, ICAM, VCAM, E-selectin, thrombin, bradykinin, PGF2 and cell surface molecules. The technique involves complexation of the target molecule with a mixture of oligonucleotides containing random sequences and sequences which serve as primer for PCR under conditions wherein a complex is formed with the specifically binding sequences, but not with the other members of the oligonucleotide mixture. The complex is then separated from uncomplexed oligonucleotides and the complexed members of the oligonucleotide mixture are recovered from the separated complex using the polymerase chain reaction. The recovered oligonucleotides may be sequenced, and successive rounds of selection using complexation, separation, amplification and recovery can be employed. The oligonucleotides can be used for therapeutic and diagnostic purposes and for generating secondary aptamers.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 20 OF 26 USPATFULL on STN

ACCESSION NUMBER:

1998:24868 USPATFULL

TITLE:

L6

Non-crosslinked protein particles for therapeutic and

diagnostic use

INVENTOR(S): PATENT ASSIGNEE(S):

Yen, Richard C. K., Yorba Linda, CA, United States Hemosphere, Inc., Irvine, CA, United States (U.S.

corporation)

NUMBER KIND DATE

PATENT INFORMATION:

19980310

APPLICATION INFO.:

us 5725804 us 1995-471650

19950606 (8)

RELATED APPLN. INFO.:

Continuation-in-part of Ser. No. US 1994-212546, filed on 14 Mar 1994, now patented, Pat. No. US 5616311 which is a continuation-in-part of Ser. No. US 1993-69831, filed on 1 Jun 1993, now abandoned And Ser. No. US 1992-959560, filed on 13 Oct 1992, now patented, Pat. No. US 5308620 which is a continuation-in-part of Ser. No. US 1991-641720, filed on 15 Jan 1991, now abandoned Utility

DOCUMENT TYPE: FILE SEGMENT:

Granted

PRIMARY EXAMINER: LEGAL REPRESENTATIVE: Lovering, Richard D. Townsend & Townsend & Crew

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

11

2178

LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Albumin particles in the nanometer and micrometer size range in an aqueous suspension are rendered stable against resolubilization without the aid of a crosslinking agent and without denaturation, by the incorporation of a stabilizing agent in the particle composition. Particles which are primarily albumin in the nanometer and micrometer size range in an aqueous suspension are rendered stable against resolublization by the incorporation of a reducing agent, oxidizing agent, hydrogen-accepting molecule, high molecular weight polymer, sulfur-containing ring compound or combinations thereof.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 21 OF 26 USPATFULL on STN L6

ACCESSION NUMBER:

97:115268 USPATFULL

TITLE: INVENTOR(S):

Synthetic catalytic free radical scavengers useful as antioxidants for prevention and therapy of disease Malfroy-Camine, Bernard, Arlington, MA, United States Doctrow, Susan Robin, Roslindale, MA, United States

PATENT ASSIGNEE(S):

Eukarion, Inc., Bedford, MA, United States (U.S.

corporation)

NUMBER KIND DATE us 5696109 us 1995-485489

PATENT INFORMATION: APPLICATION INFO.:

RELATED APPLN. INFO.:

US 1995-485489 19950607 (8)
Continuation-in-part of Ser. No. US 1995-380731, filed on 26 Jan 1995 which is a continuation-in-part of Ser. No. US 1992-987474, filed on 7 Dec 1992, now patented, Pat. No. US 5403834

NUMBER DATE

PRIORITY INFORMATION:

wo 1993-us11857 19931206

FILE SEGMENT: Granted

Jarvis, William R. A. PRIMARY EXAMINER:

LEGAL REPRESENTATIVE: Townsend and Townsend and Crew LLP

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 28 Drawing Figure(s); 19 Drawing Page(s)

LINE COUNT: 3441

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The invention provides antioxidant salen-metal complexes, compositions of such antioxidant salen-metal complexes having superoxide activity,

catalase activity, and/or peroxidase activity, compositions of salen-metal complexes in a form suitable for pharmaceutical administration to treat a disease associated with cell or tissue damage

produced by free radicals such as superoxide, and cosmetic and free radical quenching formulations of salen metal compounds.

# CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 22 OF 26 USPATFULL on STN 16

ACCESSION NUMBER: 97:26904 USPATFULL

Non-crosslinked protein particles for therapeutic and TITLE:

diagnostic use

INVENTOR(S): Yen, Richard C. K., Glendora, CA, United States PATENT ASSIGNEE(S): Hemosphere, Inc., Irvine, CA, United States (U.S.

corporation)

KIND DATE NUMBER

19970401 PATENT INFORMATION: us 5616311 US 1994-212546 APPLICATION INFO.: 19940314 (8)

RELATED APPLN. INFO.: Continuation-in-part of Ser. No. US 1993-69831, filed

on 1 Jun 1993, now abandoned And Ser. No. US 1992-959560, filed on 13 Oct 1992, now patented, Pat. No. US 5308620 which is a continuation-in-part of Ser. No. US 1991-641720, filed on 15 Jan 1991, now abandoned

DOCUMENT TYPE: Utility

FILE SEGMENT: Granted PRIMARY EXAMINER:

Lovering, Richard D.
Townsend & Townsend & Crew LEGAL REPRESENTATIVE:

NUMBER OF CLAIMS: 26 **EXEMPLARY CLAIM:** 1,26

LINE COUNT: 2585

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB Albumin particles in the nanometer and micrometer size range in an aqueous suspension are rendered stable against resolubilization without the aid of a crosslinking agent and without denaturation, by the incorporation of hemoglobin in the particle composition. Particles which are primarily hemoglobin in the nanometer and micrometer size range in an aqueous suspension are rendered stable against aggregation by the incorporation of either albumin, surface active agents or gelatin.

# CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 23 OF 26 USPATFULL ON STN

ACCESSION NUMBER: 95:29636 USPATFULL

TITLE:

Synthetic catalytic free radical scavengers useful as antioxidants for prevention and therapy of disease INVENTOR(S):

Malfroy-Camine, Bernard, Arlington, MA, United States

<--

Baudry, Michel, Long Beach, CA, United States Eukarion, Inc., Arlington, MA, United States (U.S. PATENT ASSIGNEE(S):

corporation)

NUMBER KIND DATE

PATENT INFORMATION: us 5403834 19950404 us 1992-987474 APPLICATION INFO.: 19921207 (7)

Utility DOCUMENT TYPE: FILE SEGMENT: Granted

Henley, III, Raymond Criares, T. J. PRIMARY EXAMINER: ASSISTANT EXAMINER: LEGAL REPRESENTATIVE: Dunn, Tracy J.

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 9 Drawing Figure(s); 6 Drawing Page(s)

LINE COUNT: 1742 CAS INDEXING IS AVAILABLE FOR THIS PATENT. suitable for pharmaceutical administration to treat or prevent a disease associated with cell or tissue damage produced by free radicals such as superoxide.

### CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 24 OF 26 USPATFULL on STN L6

ACCESSION NUMBER: 94:51514 USPATFULL Antiplatelet and antithrombotic activity of platelet TITLE:

glycoprotein Ib receptor fragments

INVENTOR(S): Handin, Robert, Needham, MA, United States

PATENT ASSIGNEE(S): Brigham and Women's Hospital, Boston, MA, United States

(U.S. corporation)

	NUMBER	KIND D	DATE		
PATENT INFORMATION: APPLICATION INFO.: DOCUMENT TYPE:	US 5321127 US 1991-670606 Utility		940614 910318	(7)	<
FILE SEGMENT: PRIMARY EXAMINER:	Granted Russel, Jeffrey E		0 500		

LEGAL REPRESENTATIVE: Sterne, Kessler, Goldstein & Fox

NUMBER OF CLAIMS:

EXEMPLARY CLAIM: NUMBER OF DRAWINGS:

10 Drawing Figure(s): 8 Drawing Page(s)

1494 LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

A platelet glycoprotein Ib receptor fragment, having antiplatelet and antithrombotic activity, useful for blocking platelet adhesion. The invention may be used in the treatment of patients who are particularly prone to thrombosis and embolism. The invention may also be used to purify von Willebrands factor.

## CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 25 OF 26 BIOSIS COPYRIGHT 2004 BIOLOGICAL ABSTRACTS INC. on STN \_6 1999:184941 BIOSIS ACCESSION NUMBER:

PREV199900184941

DOCUMENT NUMBER: \*\*\*cytomegalovirus\*\*\* -activated Human TITLE:

and p21Cip1 degradation in human lung fibroblasts.

Chen, Z.; Knutson, E.; Kurosky, A.; Liu, S.; Albrecht, T.

Univ. Texas Med. Branch, Galveston, TX 77555, USA CORPORATE SOURCE:

Proceedings of the American Association for Cancer Research SOURCE:

Annual Meeting, (March, 1999) Vol. 40, pp. 447-448. print. Meeting Info.: 90th Annual Meeting of the American Association for Cancer Research. Philadelphia, Pennsylvania, USA. April 10-14, 1999. American Association

for Cancer Research. ISSN: 0197-016X. Conference; (Meeting)

Conference; Abstract; (Meeting Abstract)

\_ANGUAGE: English

AUTHOR(S):

DOCUMENT TYPE:

AUTHOR(S):

ENTRY DATE:

Entered STN: 5 May 1999 Last Updated on STN: 5 May 1999

ANSWER 26 OF 26 BIOSIS COPYRIGHT 2004 BIOLOGICAL ABSTRACTS INC. on STN -6

1993:407472 BIOSIS ACCESSION NUMBER: PREV199396073197 DOCUMENT NUMBER:

Inhibition of proteolytic activity of poliovirus and TITLE:

rhinovirus 2A proteinases by elastase-specific inhibitors. Molla, Akhteruzzaman; Hellen, Christopher U. T.; Wimmer, Eckard [Reprint author]

CORPORATE SOURCE: Dep. Microbiol., Sch. Med., State Univ. New York at Stony

Brook, Stony Brook, NY 11794-8621, USA Journal of Virology, (1993) Vol. 67, No. 8, pp. 4688-4695. CODEN: JOVIAM. ISSN: 0022-538X. SOURCE:

DOCUMENT TYPE: Article \_ANGUAGE: English

ENTRY DATE: Entered STN: 8 Sep 1993

Last Updated on STN: 6 Nov 1993

A polyprotein cleavage assay has been developed to assay the proteolytic activities in vitro of the 2A proteinases encoded by poliovirus and human ۱В rhinovirus 14, which are representative members of the Enterovirus and Rhinovirus genera of picornaviruses, respectively. The elastase-specific substrate-based inhibitors elastatinal and methoxysuccinyl-Ala-Ala-Pro-Valchloromethylketone (MPCMK) inhibited both 2A proteinases in vitro. The

incubation with elastatinal, whereas the mobility of a Cys-109 fwdarw -Ala poliovirus 2A-pro mutant was unchanged, an observation suggesting that this inhibitor may have formed a covalent bond with the active-site Cys-109 nucleophile. Iodoacetamide, \*\*\*calpain\*\*\* inhibitor 1, and antipain inhibited poliovirus 2A-pro. MPCMK caused a reduction in the inhibitor 1, and yields of the enteroviruses poliovirus type 1 and coxsackievirus A21 and of human rhinovirus 2 in infected HeLa cells but did not affect the growth of encephalomyocarditis virus, a picornavirus of the Cardiovirus genus.
MPCMK abrogated the shutoff of host cell protein synthesis that is induced by enterovirus and rhinovirus infection and reduced the synthesis of virus-encoded polypeptides in infected cells. These results indicate that the determinants of substrate recognition by 2A proteinases resemble those of pancreatic and leukocyte elastases. These results may be relevant to the development of broad-range chemotherapeutic agents against entero- and rhinoviruses.

# d 16 ibib kwic tot

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ANSWER 1 OF 26 CAPLUS COPYRIGHT 2004 ACS on STN
                        1999:723196 CAPLUS
ACCESSION NUMBER:
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OCUMENT NUMBER: 131:333006

Production of recombinant replication-deficient viral TTLE:

vectors encoding exogenous transgenes via

APPLICATION NO.

DATE

microcarrier-based process

INVENTOR(S): Giroux, Daniel D.; Goudreau, Ann M.; Ramachandra,

Muralidhara; Shabram, Paul W.

Canji, Inc., USA PCT Int. Appl., 32 pp. PATENT ASSIGNEE(S): OURCE:

KIND

CODEN: PIXXD2

DATE

OCUMENT TYPE: Patent English .ANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.

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             KZ, LC, LK, LR, LT, LU, LV, MD, MG, MK, MN, MX, NO, NZ, PL, PT,
             RO, RU, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UZ, VN, YU, ZA,
             AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
        RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

1994134

A 19991130

US 1998-73076

19980504 <--
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    AU 9938823
                              19991123
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                                                                 19990504 <--
                                              EP 1999-921681
                                                                 19990504
    EP 1078095
                              20010228
                        A1
             AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, PT, IE,
    LT, LV, FI, RO
JP 2002513583 T2
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                                              JP 2000-547250
                                                                 19990504
                                           us 1998-73076
RIORITY APPLN. INFO.:
                                                                 19980504
                                           wo 1999-us9813
                                                                 19990504
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EFERENCE COUNT:
                                 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS
                                 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
                       ***19991111***
    WO 9957297 A1
    PATENT NO.
                       KIND DATE
                                              APPLICATION NO.
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                              19991111
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             AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
        RW: GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW, AT, BE, CH, CY, DE, DK,
             ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG,
             CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
34 A 19991130 US 1998-73076
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                                              CA 1999-2328084
                                                                 19990504 <--
                        AΑ
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                                              AU 1999-38823
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                              20010228
                                              EP 1999-921681
                                                                 19990504
             AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, PT, IE,
             LT, LV, FI, RO
                                              JP 2000-547250
    JP 2002513583
                        T2
                             20020514
                                                                 19990504
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RL: BAC (Biological activity or effector, except adverse); BSU (Biological
     study, unclassified); BIOL (Biological study)
(early, ***cytomegalovirus***; prodn. of recombinant
         replication-deficient viral vectors encoding exogenous transgenes via
         microcarrier-based process)
044-82-1, ***Calpain***
     110044-82-1,
                                            inhibitor I
     RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BIOL (Biological study)
         (prodn. of recombinant replication-deficient viral vectors encoding
         exogenous transgenes via microcarrier-based process)
     ANSWER 2 OF 26 CAPLUS COPYRIGHT 2004 ACS on STN
                              1999:718875
CCESSION NUMBER:
                                              CAPLUS
OCUMENT NUMBER:
                              131:348774
                              Tandem fluorescent protein constructs and their
ITLE:
                              preparation for enzyme assays
                              Tsien, Roger Y.; Heim, Roger; Cubitt, Andrew
The Regents of the University of California, USA;
NVENTOR(S):
ATENT ASSIGNEE(S):
                              Aurora Biosciences Corporation
                              U.S., 33 pp., Cont.-in-part of U.S. Ser. No. 594,575. CODEN: USXXAM
OURCE:
OCUMENT TYPE:
                              Patent
ANGUAGE:
                              English
AMILY ACC. NUM. COUNT:
ATENT INFORMATION:
     PATENT NO.
                          KIND DATE
                                                     APPLICATION NO.
                                                                           DATE
                                  19991109
                                                                           19970131 <--
     us 5981200
                                                     us 1997-792553
                                                     PT 1997-905667
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     PT 877805
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     ES 2177939
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     us 2003186229
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     us 2002164674
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                                                                       A2 19960131
RIORITY APPLN. INFO.:
                                                 us 1996-594575
                                                 us 1997-792553
                                                                       A1 19970131
                                                 us 1999-396003
                                                                       B2 19990913
                                      THERE ARE 22 CITED REFERENCES AVAILABLE FOR THIS
EFERENCE COUNT:
                              22
                                      RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
                          ***19991109***
    US 5981200 A
    PATENT NO.
                          KIND DATE
                                                     APPLICATION NO. DATE
                                                     US 1997-792553
PT 1997-905667
ES 1997-905667
US 2001-865291
US 2002-57505
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     us 2003186229
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    us 2002164674
                           Α1
                                                                           20020125
    This invention provides tandem fluorescent protein construct including a
    donor fluorescent protein moiety, an acceptor fluorescent protein moiety and a linker moiety that couples the donor and acceptor moieties. The
     donor and acceptor moieties exhibit fluorescence resonance energy transfer
    which is eliminated upon cleavage. The constructs are useful in enzymic assays. Mutant green fluorescent proteins (GFPs) were created by mutagenesis of the Aequorea victoria GFP. Polyhistidine tagged tandem green and blue fluorescent proteins were recombinantly constructed having an inserted periode sequence including cleavage recognition sites for many
                    Cleavage expts. were done with trypsin, enterokinase and
     proteases.
       ***calpain***
     78990-62-2,
                      ***Calpain***
    RL: ANT (Analyte); PEP (Physical, engineering or chemical process); ANST (Analytical study); PROC (Process)
         (fluorescent fusion protein cleavage with; tandem fluorescent protein constructs and their prepn. for enzyme assays)
     139691-88-6, Assemblin
     RL: ANT (Analyte); THU (Therapeutic use); ANST (Analytical study); BIOL
     (Biological study); USES (Uses)
(of ***cytomegalovirus***
                                              ; tandem fluorescent protein constructs
         and their prepn. for enzyme assays)
    ANSWER 3 OF 26 CAPLUS COPYRIGHT 2004 ACS on STN
                              1998:728567 CAPLUS
CCESSION NUMBER:
OCUMENT NUMBER:
                              130:10614
ITLE:
                              Ricin precursors cleavable by disease-specific
                              proteinases for treatment of cancer, viral or
                              parasitic infections
NVENTOR(S):
                              Borgford, Thor
ATENT ASSIGNEE(S):
                              De Novo Enzyme Corp., Can.
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Т

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Ι

Т

Т

6

CODEN: PIXXD2

OCUMENT TYPE: ANGUAGE: Patent English

AMILY ACC. NUM. COUNT: :

ATENT INFORMATION:

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PATENT NO.
                            KIND DATE
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                                                          us 1999-403752
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    us 2004009551
                                     20040115
                                                          us 2003-394511
                                                                                  20030324
                              Α1
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RIORITY APPLN. INFO.:
                                                                                  19970430
                                                                              Ρ
                                                                                  19971029
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                                                      WO 1998-CA394
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                                                      us 1999-403752
                                                                            A3 19991029
                            ***19981105***
    WO 9849311 A2
    PATENT NO.
                            KIND DATE
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                                                                                 DATE
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37 A1 19981124 AU 19
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    US 2004009551
                                     20040115
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                              Α1
                                                                                  20030324
         (pAP260, transfer vector for
                                                  ***calpain*** -cleavable preproricin
         gene; ricin precursors cleavable by disease-specific proteinases for
         treatment of cancer, viral or parasitic infections)
         (pAP262, transfer vector for ***calpain*** -cleavable preproricin gene; ricin precursors cleavable by disease-specific proteinases for
         treatment of cancer, viral or parasitic infections)
                                                  ***calpain*** -cleavable preproricin
         (pAP294, transfer vector for
         gene; ricin precursors cleavable by disease-specific proteinases for
    treatment of cancer, viral or parasitic infections)

152870-66-1 215649-49-3 215649-50-6 215649-51-7

RL: BPR (Biological process); BSU (Biological study, unclassified); PRP (Properties); BIOL (Biological study); PROC (Process)

( ***calpain*** -labile linker for ricin precursor; ricin precursors
         cleavable by disease-specific proteinases for treatment of cancer,
         viral or parasitic infections)
     215649-39-1
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    RL: BPR (Biological process); BSU (Biological study, unclassified); PRP (Properties); BIOL (Biological study); PROC (Process) ( ***cytomegalovirus*** proteinase-labile linker for ricin precursor; ricin precursors cleavable by disease-specific proteinases
         for treatment of cancer, viral or parasitic infections)
    215649-52-8
    RL: BPR (Biological process); BSU (Biological study, unclassified); PRP
     (Properties); BIOL (Biological study); PROC (Process)
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ricin precursors cleavable by disease-specific proteinases for
        treatment of cancer, viral or parasitic infections) 649-41-5 215649-42-6
T.
    215649-41-5
    proteinases for treatment of cancer, viral or parasitic infections) 805-06-4 215805-20-2 215861-07-7 215861-33-9
T
    RL: BSU (Biological study, unclassified); PRP (Properties); BIOL
     (Biological study)
        (nucleotide sequence,
                                    ***calpain***
                                                       cleavage of preproricin in
    relation to; ricin precursors cleavable by disease-specific proteinases for treatment of cancer, viral or parasitic infections)
215804-74-3 215804-75-4 215804-99-2
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     (Biological study)
        (nucleotide sequence, encoding ***calpain*** cleavage site; ricin precursors cleavable by disease-specific proteinases for treatment of
                                              ***calpain***
        cancer, viral or parasitic infections)
802-07-6 215802-08-7
     215802-07-6
Τ.
    RL: BUU (Biological use, unclassified); PRP (Properties); BIOL (Biological
    study); USES (Uses)

(nucleotide sequence, encoding cleavage site for proteinase of human

***cytomegalovirus*** ; ricin precursors cleavable by disease-specific
        proteinases for treatment of cancer, viral or parasitic infections)
    RL: BSU (Biological study, unclassified); PRP (Properties); BIOL
     (Biological study)
                                    ***herpes***
                                                         ***simplex***
        (nucleotide sequence,
                                                                            virus
        proteinase cleavage of preproricin in relation to; ricin precursors
        cleavable by disease-specific proteinases for treatment of cancer,
        viral or parasitic infections)
    215802-67-8
                     215802-76-9
    RL: BSU (Biological study, unclassified); PRP (Properties); BIOL
     (Biological study)
         (nucleotide sequence, human _***cytomegalovirus***
                                                                         proteinase
        cleavage of preproricin in relation to; ricin precursors cleavable by disease-specific proteinases for treatment of cancer, viral or parasitic infections)
                                                          9025-26-7, Cathepsin D
     9001-01-8, Kallikrein
                                 9004-06-2, Elastase
Т
     9039-53-6, Urokinase 9047-22-7, Cathepsin B 60616-82-2, Cathepsin L 69458-91-9, Candidapepsin 78990-62-2, ***Calpain*** 79955-99-0,
    Matrix metalloproteinase 3
                                       139691-88-6
                                                      141256-52-2, Matrix
                              141907-41-7, Matrix metalloproteinase
                                                                               146480-36-6,
    metalloproteinase 7
    Matrix metalloproteinase 9
     RL: CAT (Catalyst use); THU (Therapeutic use); BIOL (Biological study);
    USES (Uses)
         (ricin precursors cleavable by; ricin precursors cleavable by
        disease-specific proteinases for treatment of cancer, viral or
        parasitic infections)
    ANSWER 4 OF 26 USPATFULL ON STN
                           2003:296940 USPATFULL
CCESSION NUMBER:
TITLE:
                           Lactacystin analogs
                           Schreiber, Stuart L., Boston, MA, United States
Standaert, Robert F., Bryan, TX, United States
INVENTOR(S):
                           Fenteany, Gabriel, Cambridge, MA, United States
Jamison, Timothy F., Cambridge, MA, United States
                           Millennium Pharmaceuticals, Inc., Cambridge, MA, United
PATENT ASSIGNEE(S):
                           States (U.S. corporation)
                                 NUMBER
                                                KIND
                                                         DATE
PATENT INFORMATION:
                           us 6645999
                                                  в1
                                                        20031111
                           wo 9632105
                                                        19961017
                           us 1997-945092
                                                        19970126
APPLICATION INFO.:
                                                                    (8)
                           wo 1996-us5072
                                                        19960412
                                                        19980126 PCT 371 date
RELATED APPLN. INFO.:
                           Continuation-in-part of Ser. No. US 1995-421583, filed
                           on 12 Apr 1995
Utility
OCUMENT TYPE:
ILE SEGMENT:
                           GRANTED
PRIMARY EXAMINER:
                           Travers, Russell
EGAL REPRESENTATIVE:
                           Hale and Dorr LLP
NUMBER OF CLAIMS:
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NUMBER OF DRAWINGS:
                            0 Drawing Figure(s); 0 Drawing Page(s)
                            2868
LINE COUNT:
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
        us 6645999
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                                    20031111
        wo 9632105
                      19961017
        The compounds disclosed herein are highly selective for the proteasome,
SUMM
        and do not inhibit other proteases such as trypsin, .alpha.-chymotrypsin, ***calpain*** I, ***calpain*** II, pap
                                                                      II, papain, and
        cathepsin B.
SUMM
                   the X/MB1 subunit and .alpha.-chain of the proteasome and do
        not inhibit the activity of proteases such as trypsin,
        .alpha.-chymotrypsin,
                                    ***calpain***
                                                     Į,
                                                             ***calpain***
        cathepsin, and papain. Such selectivity is useful to formulate a pharmaceutical composition with fewer side effects and to evaluate.
SUMM
        Other eukaryotic transcription factors that require proteolytic
        processing include the general transcription factor TFIIA,
***herpes*** ***simplex*** virus VP16 accessory pr
                                               virus VP16 accessory protein (host cell
        factor), virus-inducible IFN regulatory factor 2 protein, and the
        DETD
        for assay); Papain: 50.
DETD
TABLE 4
Inhibition of Other Proteases
  Effect of lactacystin
 Protease tested (100 .mu.M)
 .alpha.-Chymotrypsin No inhibition
 Trypsin No inhibition
      ***Calpain***
                        I No inhibition
      ***Calpain***
                        II No inhibition
 Papain No inhibition
 Cathepsin B No inhibition
                   containing the human p105 cDNA. Forty-eight hrs after
DETD
        transfection, cells were pretreated for 1 hour with 0.5% DMSO, 50 .mu.M
           ***calpain***
                            inhibitor II, 50 .mu.M MG132, or 10 .mu.M
        .beta.-lactone. Cells were then pulse labelled with 250 uCi/plate of
        .sup.35S-methionine/cysteine for.
        apparent, as was expected (Fan and Maniatis, 1991, Nature 354:395; Palombella et al., 1994, Cell 78:773). Pretreatment of cells with ***calpain*** inhibitor II has no effect on the processing of p105 to p50 (lane 4). However, treatment of cells with the. . .
DETD
     ANSWER 5 OF 26 USPATFULL ON STN
                            2003:190684 USPATFULL
ACCESSION NUMBER:
TITLE:
                            Ricin-like toxin variants for treatment of cancer,
                            viral or parasitic infections
Borgford, Thor, Burnaby, CANADA
Twinstrand Therapeutics Inc., Vancouver, CANADA
INVENTOR(S):
PATENT ASSIGNEE(S):
                            (non-U.S. corporation)
                                  NUMBER
                                                KIND
                                                          DATE
PATENT INFORMATION:
                            us 6593132
                                                в1
                                                        20030715
                                                        19981105
                            wo 9849311
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                            US 1999-403752
                                                        19991029
APPLICATION INFO.:
                                                                    (9)
                            WO 1998-CA394
                                                        19980430
                            Utility
DOCUMENT TYPE:
FILE SEGMENT:
                            GRANTED
PRIMARY EXAMINER:
                            Carlson, Karen Cochrane
LEGAL REPRESENTATIVE:
                            Bereskin & Parr, Gravelle, Micheline
NUMBER OF CLAIMS:
                            36
EXEMPLARY CLAIM: NUMBER OF DRAWINGS:
                            254 Drawing Figure(s); 254 Drawing Page(s)
LINE COUNT:
                            5176
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
        US 6593132
PΙ
                                    20030715
                              В1
        wo 9849311 19981105
SUMM
                   (Proc. Natl. Acad. Sci. USA 88:107973-10800 (1991)) has
        described a series of viral proteases which are specifically associated
```

```
virus, varicella zoster virus-I. and infectious laryngotracheitis virus.
              These proteases possess similar substrate specificity and play an.
             . . . breast cancer, prostate cancer, non small cell lung cancer, malaria, and diverse viral disease states associated with infection with human ***cytomegalovirus***, hepatitis virus, herpes virus, human rhinovirus, infectious laryngotracheitis virus, poliomyelitis virus, or
SUMM
              varicella zoster virus.
                               SLSALLSSDIFN cleaved by human prostate-specific antigen;
SUMM
              SLPRFKIIGGFN cleaved by kallikrein (bK3): SLLGIAVPGNFN cleaved by
             neutrophil elastase; and FFKNIVTPRTPP cleaved by ***calpain***

(calcium activated neutral protease). The nucleic acid sequences for ricin A and B chains with each of the linker sequences.

. . . serine, asparagine or valine. In particular embodiments, the linker amino acid sequence comprises SGVVNASCRLAN or SSYVKASVSPEN
SUMM
                                                     `***cytomegalovirus***
                                                                                                  protease; SALVNASSAHVN or
              cleaved by a human
              STYLQASEKFKN cleaved by a ***herpes***
                                                                                                   ***simplex***
                                                                                                                                  1 virus
              protease; SSILNASVPNFN cleaved by a human herpes virus 6 protease;
              SQDVNAVEASSN or SVYLQASTGYGN cleaved by a varicella zoster.
             SQDVNAVEASSN or SVYLQASTGYGN cleaved by a varicella zoster.
. . . In a particular embodiment, the cancer is T-cell or B-cell lymphoproliferative disease. In another particular embodiment, the virus is human ***cytomegalovirus***, Epstein-Barr virus, hepatitis virus, herpes virus, human rhinovirus, infectious laryngotracheitis virus, poliomyelitis virus, or varicella zoster virus. In a further. . . . . . cell carcinoma, gastrointestinal cancer, breast cancer, prostate cancer, non small cell lung cancer. In another embodiment, the virus is human ***cytomegalovirus***, Epstein-Barr virus, hepatitis
SUMM
SUMM
              virus, herpes virus, human rhinovirus, human T-cell leukemia virus,
              infectious laryngotracheitis virus, póliomyelitis virus, or varicella
              FIG. 47B shows the nucleotide sequence of the ***calpain***
DRWD
              region of pAP-296;
             FIG. 47D shows the amino acid sequence comparison of mutant preproricin linker region of ***calpain*** to wild type;
DRWD
             FIG. 64 is a blot showing cleavage of pAP-296 with The nucleotide sequence of the ***calpain*** 1
                                                                                                                  ***calpain***
DRWD
                                                                                                          linker region of
DETD
             pAP-296 is referred to herein as SEQ ID NO. 124.
             The amino acid sequence of the mutant preproricin linker region for ***calpain*** , pAP-296, is referred to herein as SEQ ID NO. 126.

In a further embodiment, the preparation of proteases from human ***cytomegalovirus*** , human herpes virus, varicalla zoster virus and
DETD
DETD
             infectious laryngotracheitis virus have been taught by Liu F. & Roizman,
DETD
                              for directing expression in mammalian cells generally include a
             promoter (e.g., derived from viral material such as polyoma, Adenovirus
             2, ***cytomegalovirus*** and Simian Virus 40), as well as other transcriptional and translational control sequences. Examples of
             mammalian expression vectors include pCDM8.
                                                                                                  ***Cytomegalovirus***
DETD
             Cleavage of pAP-248 Protein with the Human
              (HCMV) Protease
DETD
                                                                               ***Calpain***
             Cleavage of pAP-296 Protein with
             . . . disease-specific proteases to confirm specific cleavage in the linker region. The pAP-296 protein sample (2.05 ug) was digested with the ***Calpain*** protease (10 ug) overnight at 37.degree. C. The total volume of the digestion reaction was 35 ul and 0.761 ug of the reaction sample was loaded on a protein gel. The ***Calpain***
DETD
             protease was purchased from Sigma Chemical Co., USA . . . 58 & 66(MMP-2), 60, 64 and 62 show the cleavage of proteases of
DETD
             linkers by HCMV, HAV 3C, MMP-2, t-PA,
                                                                                          ***calpain*** , and human
             neutraphil elastase respectively. Without protease digestion, the proricin variants appear as a single band at approximately 60 kDA. . . . HCMV; pAP-256 by HAV3C protease; pAP-270 by MMP-2 protease; pAP-288 by t-PA protease; pAP-294 by human neutrophil elastase; pAP-296 by ***calpain***; and pAP-222 by MMP-2 is illustrated in FIGS. 52, 59, 61, 63, 65, and 67 respectively. The appearance of. . .
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                             Val Pro Gly Asn Phe Asn
DETD
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<212> TYPE: DNA
<213> ORGANISM:
                                  ***Calpain***
                                                                linker region of pAP-296
<400> SEQUENCE: 124
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<210> SEQ ID NO 125
 <211> LENGTH: 1855
 <212>. . agcaagttat atcgaattcc tgcag
                                                                                                    1855
 <210> SEQ ID NO 126
 <211> LENGTH: 12
 <212> TYPE: PRT
 <213> ORGANISM: Mutant preproricin linker region for ***calpain*** , pAP-296
 <400> SEQUENCE: 126
 Phe Phe Lys Asn Ile Val Thr Pro Arg Thr Pro Pro
 CLM
               What is claimed is:
                     a matrix metalloproteinase, cathepsin L, cathepsin D, urokinase-type
               plasminogen activator, tissue-type plasminogen activator, human
prostate-specific antigen, kallikrein, neutrophil elastase, and
                   ***calpain***
                     a cleavage recognition site for a viral protease, wherein the viral
               protease is selected from the group consisting of: human
                   ***cytomegalovirus***
                                                                  , human herpes virus, varicella zoster virus,
               hepatitis A virus, hepatitis C virus, Epstein-Barr virus-specific
               protease, and infectious laryngotracheitis virus.
          ANSWER 6 OF 26 USPATFULL on STN
ACCESSION NUMBER:
                                                 2002:115819 USPATFULL
TITLE:
                                                 Fibrinogen-coated particles for therapeutic use
INVENTOR(S):
                                                 Yen, Richard C. K., Yorba Linda, CA, United States
                                                 Hemosphere, Inc., Anaheim, CA, United States (U.S.
PATENT ASSIGNEE(S):
                                                 corporation)
                                                           NUMBER
                                                                                     KIND DATE
PATENT INFORMATION:
                                                 US 6391343
                                                                                                  20020521
                                                 wo 9639128
                                                                                                  19961212
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                                                 US 1998-952765
APPLICATION INFO.:
                                                                                                  19980410
                                                 WO 1996-US9458
                                                                                                  19960604
                                                 19980410 PCT 371 date
Continuation-in-part of Ser. No. US 1995-554919, filed on 9 Nov 1995, now abandoned Continuation-in-part of
RELATED APPLN. INFO.:
                                                 Ser. No. US 1995-471650, filed on 6 Jun 1995, now
                                                 patented, Pat. No. US 5725804 Continuation-in-part of
                                                 Ser. No. US 1994-212546, filed on 14 Mar 1994, now
                                                 patented, Pat. No. US 5616311 Continuation-in-part of
                                                 Ser. No. US 1993-69831, filed on 1 Jun 1993, now abandoned Continuation-in-part of Ser. No. US 1992-959560, filed on 13 Oct 1992, now patented, Pat. No. US 5308620 Continuation-in-part of Ser. No. US 1993-698620 Continuation-part of Ser. No. US 1993-698620 Continuation-in-part of Ser. No. US 1993-69881, filed on 1 Jun 1993, now abandoned Continuation-in-part of Ser. No. US 1993-69881, filed on 1 Jun 1993, now abandoned Continuation-in-part of Ser. No. US 1993-69881, filed on 1 Jun 1993, now abandoned Continuation-in-part of Ser. No. US 1993-69881, filed on 1 Jun 1993, now abandoned Continuation-in-part of Ser. No. US 1993-95960, filed on 13 Oct 1992, now patented, Pat. No. US 1993-69881, patented in paten
                                                 1991-641720, filed on 15 Jan 1991, now abandoned
DOCUMENT TYPE:
                                                 Utility
FILE SEGMENT:
                                                 GRANTED
PRIMARY EXAMINER:
                                                 Lovering, Richard D.
LEGAL REPRESENTATIVE:
                                                 Townsend and Townsend and Crew LLP
NUMBER OF CLAIMS:
EXEMPLARY CLAIM:
NUMBER OF DRAWINGS:
                                                 0 Drawing Figure(s); 0 Drawing Page(s)
LINE COUNT:
                                                 2407
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
PΙ
              US 6391343
                                                     в1
                                                               20020521
                                      19961212
              wo 9639128
                                                                                                                                           <--
DETD
                   ***Calpain***
                                                   Inhibitor I
                   ***Calpain***
DETD
                                                   Inhibitor II
                   ***Calpain***
DETD
                                                   Inhibitor Peptide
              DNA, M13 DNA, Adenovirus DNA, phi-X 174 phage DNA, Simian virus DNA, phi-X 174 phage DNA, Simian virus DNA,
DETD
                   ***cytomegalovirus*** DNA, Epstein-Barr Virus genes, ***Herpes***
```

genes, ribosomal RNA, human DNA and RNA; Genes coding

for ribozymes; genes coding for antibiotics (e.g., ampicillin,

chloramphenicol, cycloserine, gentamycin,.

\*\*\*Simplex\*\*\*

```
CESSION NUMBER:
                          2001:82299 USPATFULL
                          Method and product for cleaning and/or whitening of
TLE:
                          teeth
IVENTOR(S):
                          Rinne, Ari, Pajutie 3 B, FIN-2G900 Turku, Finland
                                NUMBER
                                                KIND
                                                         DATE
TENT INFORMATION:
                          US 6241973
                                                       20010605
                          wo 9829088
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                          us 1999-331777
PLICATION INFO.:
                                                       19990624
                                                                   (9)
                          WO 1998-FI1
                                                       19980102
                                                       19990624
                                                                   PCT 371 date
                                                                  PCT 102(e) date
                                                       19990624
                                  NUMBER
                                                   DATE
IORITY INFORMATION:
                          FI 1997-12
                                                 19970103
CUMENT TYPE:
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LE SEGMENT:
                          Granted
IMARY EXAMINER:
                          Rose, Shep K.
GAL REPRESENTATIVE:
                          Lydon, James C.
IMBER OF CLAIMS:
EMPLARY CLAIM:
NE COUNT:
                          583
S INDEXING IS AVAILABLE FOR THIS PATENT.
      US 6241973
                            В1
                                  20010605
      wo 9829088 19980709
MM
      Among the mammalian cysteine proteinases are further known
     calcium-activated cysteine proteinases, which are considered to belong to the ***calpain*** family. Their inhibitors are called calpastatins (M. Nakamura, s. Imajoh-Ohmi, K, Suzuki and S. Kawashima: An endogenous inhibitor of calcium-activated. . .
                 3. Installment 1995; Bjorck, L., Grubb, A. and Kjellen, L.
MM
      (1990) Cystatin C, a human proteinase inhibitor, blocks replication of ***Herpes*** ***simplex*** virus. J Virol 64, 941-943Bjorck, L
                                              virus. J Virol 64, 941-943Bjorck, L.,
     Akesson, P., Bohus, M., Trojnar, J., Abrahamson, M., Olafson, I., and Grubb, A. (1989). . . .
                protein chemistry as well as produced by molecular biological
MM
      techniques. Most of such cysteine proteinases belong to the cathepsin or
        ***calpain***
                           family.
   ANSWER 8 OF 26 USPATFULL on STN
CESSION NUMBER:
                          2000:41033
                                       USPATFULL
TLE:
                          Synthetic catalytic free radical scavengers useful as
                          antioxidants for prevention and therapy of disease
Malfroy-Camine, Bernard, Arlington, MA, United States
VENTOR(S):
                          Doctrow, Susan Robin, Roslindale, MA, United States
                          Eukarion, Inc., Bedford, MA, United States (U.S.
TENT ASSIGNEE(S):
                          corporation)
                                NUMBER
                                               KIND
                                                         DATE
                          US 6046188
TENT INFORMATION:
                                                       20000404
                          wo 9640148
                                                       19961219
                                                                               <--
                          us 1998-973577
                                                       19980311
PLICATION INFO.:
                                                                   (8)
                                                       19960606
                          wo 1996-us10037
                                                       19980311
                                                                  PCT 371 date
                                                       19980311
                                                                  PCT 102(e) date
                          Continuation-in-part of Ser. No. US 1995-485489, filed
LATED APPLN. INFO.:
                          on 7 Jun 1995, now patented, Pat. No. US 5696109
CUMENT TYPE:
                          Utility
LE SEGMENT:
                          Granted
IMARY EXAMINER:
                          Reamer, James H.
                          Townsend & Townsend & Crew LLP
GAL REPRESENTATIVE:
MBER OF CLAIMS:
                          24
EMPLARY CLAIM:
MBER OF DRAWINGS:
                          28 Drawing Figure(s); 16 Drawing Page(s)
                          3405
NE COUNT:
S INDEXING IS AVAILABLE FOR THIS PATENT.
     US 6046188
                                  20000404
     wo 9640148
                  19961219
     disease, gastric ulcers, oxygen toxicity, burned patients, renal failure attendant to transplantation, and ***herpes*** ***simplex***
MM
MM
                (3) one or more oxyradical inhibitors, such as desferrioxamine
```

```
***calpain***
                                inhibitors. The formulations of these compositions is
     dependent upon the specific pathological condition sought to be treated
     or prevented, the.
                    active ingredients, typically selected from the group
     consisting of: N-2-mercaptopropionylglycine, N-acetylcysteine, glutathione, dimethyl thiourea, desferrioxamine, mannitol, .alpha.-tocopherol, ascorbate, allopurinol, 21-aminosteroids, ***calpain*** inhibitors, glutamate receptor antagonists, plasminogen activator, streptokinase, urokinase, nonsteroidal
     anti-inflammatory agent, cortisone, and carotenoids. Antioxidant
     salen-Mn complexes may also.
  ANSWER 9 OF 26 USPATFULL on STN
                                 1999:166974 USPATFULL
ESSION NUMBER:
LE:
                                 Cysteine protease inhibitors
                                 Spruce, Lyle W., Chula Vista, CA, United States
ENTOR(S):
                                 Gyorkos, Albert C., Westminster, CO, United States
                                 Cheronis, John C., Conifer, CO, United States
Goodfellow, Val S., Tucson, AZ, United States
                                Leimer, Axel H., Westborough, MA, United States
Young, John M., Redwood City, CA, United States
Gerrity, James Ivan, Albany, OR, United States
Cortech Inc., Bedminster, NJ, United States
Corporation)
ENT ASSIGNEE(S):
                                         NUMBER
                                                              KIND
                                                                           DATE
ENT INFORMATION:
                                 US 6004933
                                                                        19991221
                                                                                                         <--
LICATION INFO.:
                                 US 1998-65258
                                                                        19980423 (9)
                                            NUMBER
                                                                  DATE
ORITY INFORMATION:
                                 US 1997-44819P
                                                             19970425 (60)
JMENT TYPE:
                                 Utility
E SEGMENT:
                                 Granted
                                Richter, Johann
Solola, Taofiq A.
Dechert Price & Rhoads
MARY EXAMINER:
ISTANT EXAMINER:
AL REPRESENTATIVE:
BER OF CLAIMS:
                                 146
MPLARY CLAIM:
BER OF DRAWINGS:
                                 4 Drawing Figure(s); 3 Drawing Page(s)
E COUNT:
                                 2591
INDEXING IS AVAILABLE FOR THIS PATENT.
                                            19991221
     US 6004933
     . . . comprise a family of intracellular cysteine proteases which are ubiquitously expressed in mammalian tissues. Three major calpains have been identified: ***calpain*** I and II, and p94. The ***calpain*** family of cysteine proteases has been implicated in many diseases and disorders, including stroke, neurodegeneration, such as
     Alzheimer's disease, amyotrophy and motor neuron damage; acute central nervous system injury, muscular dystrophy, bone resorption, platelet aggregation, cataracts and inflammation. ***Calpain*** I has been
     implicated in excitatory amino-acid induced neurotoxicity disorders
     including ischemia, hypoglycemia and epilepsy. The cysteine protease p94, a muscle-specific member of the ***calpain*** family, has been identified as a gene product responsible for limb girdle muscular
     dystrophy (Barrett A. J., et al. ICOP. where Z is a ***calpain*** binding
                                                      binding moiety, preferably R.sub.2 is
     benzyl optionally substituted with alkoxy; H.sub.2 NC(=.sup.+ NH.sub.2)NHCH.sub.2 CH.sub.2 CH.sub.2 --; --R'--C(=.sup.30
     NH.sub.2)NH.sub.2; --R'--NHC(=.sup.+.
teine proteases and exemplary recognition elements.
teine
tease P1
                                     Other
                                                    Reference
***Calpain***
                          I and II
        large hydrophobic
                      Leu, bulky
                                                    18
        e.g. Nva, Phe, Abu
                       aliphatic, hPhe
***Calpain***
        Arg or Arg-
                       t-butyl-Gly, Leu, 38
        mimetic, Lys, Tyr,
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Bzl), Leu, Abu, Phe
ain
       hPhe,.
    . . . . alpha., .beta. and .gamma.) (rice), bromelain (including stem-and fruit bromelain), ficin, thaumatopain (Thaumatococcus); gingipain R and gingipain K; calpains, including ***calpain*** (Schistosoma), ***calpain*** I, ***calpain*** II,
     (Schistosoma),
***calpain***
                            p94, calcium-binding protein PMP41, sol gene product
     (Drosophila); streptopain and cysteine endopeptidase (Porphyromonas);
     picomain 2A, picornain 3C, apothovirus endopeptidase, cardiovirus.
    sepsis), HIVinfection and AIDS, genital herpes, Zoster, chickenpox, EBV infections and encephalitis, CMV-choreoretinitis or encephalitis, ***cytomegalovirus*** infections in neonates (including related pneumonitis), opportunistic infections in immunocompromised individuals
     (including AIDS and transplant patients), dysentery, hepatitis C,
     hepatitis.
                                                             colon, kidney; osteo-,
                                   (1990); Gordon, Seminars in
            chondro-, and liposarcoma;
                                   Thrombosis and Hemostasis,
             neuroblastoma; melanoma;
                                   18:424-433 (1992)
             nonlympocytic leukemia;
             lymphocytic leukemia)
 ***Calpain***
                      I and II
            Osteoporosis, stroke, CNS
            Karlsson, et al., Neurobiology injury, Alzheimer's disease of Aging, 16:901-906 (1995);
            Additionally, diseases involving
                                   Squier, et al.,
                                                       J. Cell.
             dysregulated apoptosis as listed
                                   Physiol., 159:229-237
             for caspase above.
                                    (1994).
 ***Calpain***
                      p94 Muscular dystrophy
                                        ***Calpain***
                                                             p94 and limb-girdle
                                   muscular dystrophy, ICOP
                                   Letters 1996.
atitis C Virus
                                   Grakoui, et al., Proc. Nat.
            Hepatitis C
opeptidase 2 and
                                   Acad..
    What is claimed is:
     15. An inhibitor of claim 10 wherein Z is a
                                                                ***calpain***
                                                                                      binding
     moiety.
     109. A method of inhibiting the enzymatic activity of a
     cysteine protease comprising contacting a protease with an inhibitory
     amount of a compound of claim 15.
  ANSWER 10 OF 26 USPATFULL ON STN SION NUMBER: 1999:155518 USPATFULL
ESSION NUMBER:
                           Viral production process
LE:
                           Giroux, Daniel D., La Jolla, CA, United States
ENTOR(S):
                           Goudreau, Ann M., San Diego, CA, United States
Ramachandra, Muralidhara, San Diego, CA, United States
                           Shabram, Paul W., Olivenhain, CA, United States
Canji, Inc., San Diego, CA, United States (U.S.
ENT ASSIGNEE(S):
                           corporation)
                                 NUMBER
                                                   KIND
                                                             DATE
                           US 5994134
                                                           19991130
ENT INFORMATION:
                                                                        (9)
LICATION INFO.:
                           US 1998-73076
                                                           19980504
                           Utility
UMENT TYPE:
E SEGMENT:
                           Granted
                           Stucker, Jeffrey
MARY EXAMINER:
AL REPRESENTATIVE:
                           Murphy, Richard B.
BER OF CLAIMS:
                           17
MPLARY CLAIM:
BER OF DRAWINGS:
                           3 Drawing Figure(s); 3 Drawing Page(s)
E COUNT:
 INDEXING IS AVAILABLE FOR THIS PATENT.
    US 5994134
                                    19991130
                                                                                     <--
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val, Nle, Tyr(0-

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vectors containing a DNA sequence encoding p53 under control of the ***cytomegalovirus*** promoter region and the tripartite leader
        sequence having E3 under control of the CMV promoter and deletion of E4
        coding.
                     to the serum free media which down-regulate or inhibit the
        transgene promoter. For example, when the transgene promoter is the ***cytomegalovirus*** early promoter (CMV), elements such as
        neuramidase or tunicamycin may be added to suppress the CMV promoter
        during the culture.
                     transgene expression but should not materially interfere with
        the expression of viral genes essential to viral replication. For example, the ***cytomegalovirus*** major immediate early promoter is a promoter commonly used to constitutively drive transgene expression. This promoter contains binding sites for. . .
                    invention provides a method to increase the infectivity of
        producer cell lines for viral infectivity by the inclusion of a ***calpain*** inhibitor. Examples of ***calpain*** inhibitor.
                                                                                            inhibitors
        useful in the practice of the present invention include
                                                                                           ***calpain***
        inhibitor 1 (also known as N-acetyl-leucyl-leucyl-norleucinal, commercially available from Boehringer Mannheim). ***Calpain***
inhibitor 1 was observed to increase the infectivity of producer cell
        lines to recombinant adenovirus.
     ANSWER 11 OF 26 USPATFULL ON STN
CCESSION NUMBER:
                                1999:137014 USPATFULL
                                Vesicle transport related protein
                                Lal, Preeti, Santa Clara, CA, United States
Corley, Neil C., Mountain View, CA, United States
Shah, Purvi, Sunnyvale, CA, United States
Incyte Pharmaceuticals, Inc., Palo Alto, CA, United
NVENTOR(S):
ATENT ASSIGNEE(S):
                                States (U.S. corporation)
                                      NUMBER
                                                        KIND
                                                                  DATE
                                us 5976865
                                                                   19991102
ATENT INFORMATION:
                                US 1997-984172
                                                                   19971203 (8)
PPLICATION INFO.:
                                Utility
OCUMENT TYPE:
ILE SEGMENT:
                                Granted
                                Johnson, Nancy A.
RIMARY EXAMINER:
EGAL REPRESENTATIVE:
                                Incyte Pharmaceuticals, Inc.
UMBER OF CLAIMS:
XEMPLARY CLAIM:
                                7 Drawing Figure(s); 7 Drawing Page(s)
UMBER OF DRAWINGS:
INE COUNT:
                                2242
AS INDEXING IS AVAILABLE FOR THIS PATENT.
                                          19991102
        US 5976865
        . . . number of selection systems may be used to recover transformed cell lines. These include, but are not limited to, the ***herpes***
        ***simplex*** virus thymidine kinase genes (Wigler, M. et al. (1977) Cell 11:223-32) and adenine phosphoribosyltransferase genes (Lowy, I. et
        al. (1980).
                     a modifier of the spectrin-binding activity of ankyrin. In
        particular, cleavage of the ANK1 region from mouse erythrocyte ankyrin by ***calpain*** reduces spectrin-binding activity of the remaining 195 kDa fragment.about.8-fold (Hall, T. G. and V. Bennett (1987) J. Biol. Chem. 262:10537-45). VTRP activity is therefore measured by
        comparing the spectrin-binding activity of the 195 kDa ***calpain***
        fragment with a chimeric recombinant protein containing VTRP integrated with the 195 kDa ***calpain*** fragment. Spectrin binding is
       with the 195 kDa
       measured by incubating the VTRP recombinant protein or 195 kDa fragment with radiolabeled-.sup.14 C-spectrin together in. . .
    ANSWER 12 OF 26 USPATFULL on STN
CCESSION NUMBER:
                                1999:102423 USPATFULL
                                Method for making non-crosslinked protein particles for
                                therapeutic and diagnostic use
NVENTOR(S):
                                Yen, Richard C. K., Glendora, CA, United States
                                Hemosphere, Inc., Irvine, CA, United States (U.S.
ATENT ASSIGNEE(S):
                                corporation)
                                       NUMBER KIND
                                                                  DATE
ATENT INFORMATION:
                                US 5945033
                                                                  19990831
PPLICATION INFO.:
                                US 1996-747137
                                                                  19961112 (8)
                                Continuation of Ser. No. US 1994-212546, filed on 14
ELATED APPLN. INFO.:
                                Mar 1994, now patented, Pat. No. US 5616311 which is a
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ETD

ETD

ETD

ITLE:

ETD

ETD

ITLE:

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on 1 Jun 1993, now abandoned And Ser. No. US 1992-959560, filed on 13 Oct 1992, now patented, Pat. No. US 5308620 which is a continuation-in-part of Ser. No. US 1991-641720, filed on 15 Jan 1991, now abandoned
OCUMENT TYPE:
                                 Utility
ILE SEGMENT:
                                 Granted
                                 Dees, Jose' G.
RIMARY EXAMINER:
                                Hartley, Michael G.
Townsend and Townsend and Crew LLP
SSISTANT EXAMINER:
EGAL REPRESENTATIVE:
UMBER OF CLAIMS:
                                 12
XEMPLARY CLAIM:
                                 3655
INE COUNT:
AS INDEXING IS AVAILABLE FOR THIS PATENT.
        US 5945033
                                           19990831
                                                                                                 <--
           ***Calpain***
                                  Inhibitor I
ETD
          ***Calpain*** Inhibitor II
***Calpain*** Inhibitor Peptide
ETD
ETD
        . . . double stranded), cloning vectors, coliphage DNA, lambda phage DNA, M13 DNA, Adenovirus DNA, phi-X 174 phage DNA, Simian virus DNA, ***cytomegalovirus*** DNA, Epstein-Barr Virus genes, ***Herpes*** ***Simplex*** genes, ribosomal RNA, human DNA and RNA; Genes coding
ETD
        for ribozymes; genes coding for antibiotics (e.g., ampicillin,
        chloramphenicol, cycloserine, gentamycin,
     ANSWER 13 OF 26 USPATFULL ON STN
                                 1999:81758 USPATFULL
CCESSION NUMBER:
                                 Non-activated receptor complex proteins and uses
ITLE:
                                 Davis, Roger J., Princeton, MA, United States
NVENTOR(S):
                                 Galcheva-Gargova, Zoya, Worcester, MA, United States
ATENT ASSIGNEE(S):
                                University of Massachusetts, Boston, MA, United States
                                 (U.S. corporation)
                                        NUMBER KIND DATE .
                                us 5925566
us 1997-870518
                                                                    19990720
ATENT INFORMATION:
                                                                                                 <--
PPLICATION INFO.:
                                                                    19970606 (8)
                                          NUMBER DATE
RIORITY INFORMATION:
                                 US 1996-19219P 19960606 (60)
OCUMENT TYPE:
                                 Utility
ILE SEGMENT:
                                 Granted
                                Campell, Bruce R.
Nguyen, Dave Trong
RIMARY EXAMINER:
SSISTANT EXAMINER:
                                 Fish & Richardson, P.C.
EGAL REPRESENTATIVE:
UMBER OF CLAIMS:
                                 23
XEMPLARY CLAIM:
UMBER OF DRAWINGS:
                                 22 Drawing Figure(s); 18 Drawing Page(s)
                                 2438
INE COUNT:
AS INDEXING IS AVAILABLE FOR THIS PATENT.
                                           19990720
        US 5925566
        . . . interaction. In initial experiments, we examined the effect of proteolytic cleavage of the COOH terminus of the EGF receptor with ***calpain*** as follows. ***Calpain*** cleavage of the EGF
ETD
        receptors was performed by harvesting cells in lysis buffer without
        EDTA, PMSF, leupeptin, and aprotinin [Gregoriou. . . at 4.degree. C.
        and standard binding assays were performed as described herein. We found that both the wild-type and the ***calpain*** -cleaved EGF receptor
        that both the wild-type and the ***calpain*** -cleaved EGF receptor bound to the ZPR1 zinc fingers.

Since the major sites of ***calpain*** -cleavage of the EGF receptor are Gln.sup.996 and Asp.sup.1059 [Gregoriou et al., Eur. J. Biochem.
ETD
        223, 455 (1994)], we concluded that.
        For gene therapy, ZPR1 cDNA expression is directed from any suitable promoter (e.g., the human ***cytomegalovirus***, simian virus 40, or metallothionelip promoters), and its production is regulated by any
ETD
        desired mammalian regulatory element. For example, if. .
     ANSWER 14 OF 26 USPATFULL on STN
CCESSION NUMBER:
                                 1999:36949 USPATFULL
                                 Engineering oral tissues
ITLE:
NVENTOR(S):
                                 Mooney, David J., Ann Arbor, MI, United States
                                Rutherford, Robert B., Ann Arbor, MI, United States
The Regents of the University of Michigan, Ann Arbor,
ATENT ASSIGNEE(S):
                                 MI, United States (U.S. corporation)
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US 5885829
US 1997-864494
                                                              19990323
TENT INFORMATION:
PLICATION INFO.:
                                                               19970528
                                                                           (8)
                                       NUMBER DATE
                              US 1996-18450P 19960528 (60)
IORITY INFORMATION:
CUMENT TYPE:
                              Utility
LE SEGMENT:
                              Granted
IMARY EXAMINER:
                              Degen, Nancy
                              Arnold, White & Durkee
GAL REPRESENTATIVE:
IMBER OF CLAIMS:
                              109
EMPLARY CLAIM:
IMBER OF DRAWINGS:
                              17 Drawing Figure(s); 11 Drawing Page(s)
NE COUNT:
                              8001
S INDEXING IS AVAILABLE FOR THIS PATENT.
      US 5885829
                                       19990323
                   shown to prevent apoptosis and cell cycle effects induced by
.TD
       camptothecin and topotecan in HL-60 cells (Traganos et al., 1993).
                              inhibitor I inhibits apoptosis in thymocytes and

while leupeptin, ***calpain***
         ***Calpain***
      metamyelocytes (Squier et al., 1994), while leupeptin, ***calpain*** inhibitor II and the E64 class of serine protease inhibitors have also been shown to inhibit activation-induced programmed cell death. . .
       . . . cleavage in thymocytes without the involvement of endonucleases (Cain et al., 1994). The cysteine protease inhibitors E64 and leupeptin, the ***calpain*** selective inhibitor acetyl-leucyl-leucyl-
.TD
                                    selective inhibitor acetyl-leucyl-leucyl-
       normethional, and the serine protease inhibitors
      diisopropylfluorophosphate and phenylmethylsulfonyl fluoride were all shown to selectively block T-cell receptor-triggered. . . In various other embodiments, the human ***cytomegalovirus*** (CM immediate early gene promoter, the SV40 early promoter and the Rous
TD
       sarcoma virus long terminal repeat can be used.
                    Berkhout et al., 1988; Rowen et al., 1988; Berkhout et al., 1989; Laspia et al., 1989; Sharp and Marciniak, 1989;
TD
  Braddock et al., 1989
***Cytomegalovirus*** Weber et
                    virus*** Weber et al., 1984; Boshart et al., 1985; Foecking and Hofstetter, 1986
bbon Ape Leukemia Virus
      TD
      Human ***Cytomegalovirus*** ," Cell, 41:521, 1985.
Squier, M. K., Miller, A. C., Malkinson, A. M. and Cohen, J. J., "
***Calpain*** activation in apoptosis," J. Cell. Physiol. 159:229-237,
TD
       1994.
    ANSWER 15 OF 26 USPATFULL on STN
CESSION NUMBER:
                             1998:138941 USPATFULL
TLE:
                              Synthetic catalytic free radical scavengers useful as
                             antioxidants for prevention and therapy of disease
                             Malfroy-Camine, Bernard, Arlington, MA, United States
VENTOR(S):
                             Doctrow, Susan Robin, Roslindale, MA, United States
TENT ASSIGNEE(S):
                             Eukarion, Inc., Bedford, MA, United States (U.S.
                              corporation)
                                    NUMBER KIND DATE
TENT INFORMATION:
                             US 5834509
                                                              19981110
                             us 1995-479697
                                                              19950607
PLICATION INFO.:
                                                                           (8)
                             Continuation-in-part of Ser. No. US 1995-380731, filed on 26 Jan 1995 which is a continuation-in-part of Ser. No. US 1992-987474, filed on 7 Dec 1992, now patented,
LATED APPLN. INFO.:
                             Pat. No. US 5403834
                                      NUMBER DATE
IORITY INFORMATION:
                             wo 1993-us11857 19931206
CUMENT TYPE:
                             Utility
LE SEGMENT:
                             Granted
IMARY EXAMINER:
                             Jarvis, William R. A.
GAL REPRESENTATIVE:
                             Townsend and Townsend and Crew LLP
MBER OF CLAIMS:
EMPLARY CLAIM:
MBER OF DRAWINGS:
                             28 Drawing Figure(s); 19 Drawing Page(s)
```

NE COUNT:

3384

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. . effects in treating systemic lupus erythematosus, Crohn's
          disease, gastric ulcers, oxygen toxicity, burned patients, renal failure attendant to transplantation, and ***herpes*** ***simplex***
                            3) one or more oxyradical inhibitors, such as desferrioxamine
          or allopurinol, and/or one or more biological modifier agents, such as 
***calpain*** inhibitors. The formulations of these compositions is
                                         inhibitors. The formulations of these compositions is
          dependent upon the specific pathological condition sought to be treated
          or prevented, the.
         or prevented, the.
. . . active ingredients, typically selected from the group consisting of: N-2-mercaptopropionylglycine, N-acetylcysteine, glutathione, dimethyl thiourea, desferrioxamine, mannitol, alpha.-tocopherol, ascorbate, allopurinol, 21-aminosteroids, ***calpain*** inhibitors, glutamate receptor antagonists, tissue plasminogen activator, streptokinase, urokinase, nonsteroidal
          anti-inflammatory agent, cortisone, and carotenoids. Antioxidant
          salen-Mn complexes may also.
      ANSWER 16 OF 26 USPATFULL ON STN SSION NUMBER: 1998:134999 USPATFULL
CCESSION NUMBER:
                                         Methods for the treatment of bone resorption disorders,
ITLE:
                                         including osteoporosis
NVENTOR(S):
                                         Gelb, Bruce D., Dobbs Ferry, NY, United States
                                         Chapman, Harold, Newton, MA, United States
                                         Desnick, Robert J., New York, NY, United States
Mount Sinai School of Medicine of the City of New York,
ATENT ASSIGNEE(S):
                                         New York, NY, United States (U.S. corporation)
Brigham and Women's Hospital, Boston, MA, United States
                                          (U.S. corporation)
                                                  NUMBER
                                                                        KIND
                                                                                       DATE
                                         US 1996-704473
Utility
ATENT INFORMATION:
                                                                                      19981103
PPLICATION INFO.:
                                                                                      19960828 (8)
OCUMENT TYPE:
ILE SEGMENT:
                                         Granted
                                         Marschel, Ardin H.
Pennie & Edmonds LLP
RIMARY EXAMINER:
EGAL REPRESENTATIVE:
JMBER OF CLAIMS:
XEMPLARY CLAIM:
JMBER OF DRAWINGS:
                                         7 Drawing Figure(s); 6 Drawing Page(s)
INE COUNT:
                                         2434
AS INDEXING IS AVAILABLE FOR THIS PATENT.
                                                      19981103
         us 5830850
         . . . Science 244:1288-1292; Capecchi, 1989, Trends in Genet. 5:70-76). Utilizing the PNS method, nonhomologous recombinants are selected against by using the ***Herpes*** ***Simplex*** virus thymidine kinase (HSV-TK) gene and selecting against its PN inconting with hornor drugs such as garcyclovin on ETAL PN
         insertion with herpes drugs such as gancyclovir or FIAU. By.
         is notable that another genetic disorder (limb-girdle muscular dystrophy Type 2A) caused by the deficiency of a neutral cysteine protease, ***calpain*** 3, was recently identified in which presumably related families on a small island had different mutations (Richard, I et al., 1995, Cell 81:27-40). ***Calpain*** 3 belongs to a family of calpains, analogous to the cathepsin family. The finding of multiple mutations in ***Calpain*** 3 suggested to Richard et al. that a "digenic" model in which only in the presence of specific alleles
                                                                                                                         3 belongs to
          that a "digenic" model in which only in the presence of specific alleles
         at a permissive second locus (e.g., a compensatory, partially redundant, regulatory, or modifier gene) will there be expression of ***calpain*** mutations. Since one would need mutations at both loci to be affected, the disease prevalence would remain low. By analogy,. .
      ANSWER 17 OF 26 USPATFULL on STN
CCESSION NUMBER:
                                         1998:131743 USPATFULL
                                         Synthetic catalytic free radical scavengers useful as
                                         antioxidants for prevention and therapy of disease Malfroy-Camine, Bernard, Arlington, MA, United States Doctrow, Susan Robin, Roslindale, MA, United States Eukarion, Inc., Bedford, MA, United States (U.S.
NVENTOR(S):
ATENT ASSIGNEE(S):
                                         corporation)
                                                  NUMBER
                                                                       KIND DATE
ATENT INFORMATION:
                                         us 5827880
                                                                                     19981027
                                                                                                                           <--
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19981110

US 5834509

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**UMM** 

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ETD

ETD

[TLE:

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Continuation-in-part of Ser. No. US 1992-987474, filed
LATED APPLN. INFO.:
                          on 7 Dec 1992, now patented, Pat. No. US 5403834
                          Utility
CUMENT TYPE:
LE SEGMENT:
                          Granted
                          Nazario-Gonzalez, Porfirio
IMARY EXAMINER:
                          Townsend and Townsend and Crew LLP
GAL REPRESENTATIVE:
                          13
MBER OF CLAIMS:
EMPLARY CLAIM:
MBER OF DRAWINGS:
                          28 Drawing Figure(s); 19 Drawing Page(s)
NE COUNT:
                          3241
S INDEXING IS AVAILABLE FOR THIS PATENT.
                                  19981027
     us 5827880
     . . . effects in treating systemic lupus erythematosus, Crohn's disease, gastric ulcers, oxygen toxicity, burned patients, renal failure
MM
     attendant to transplantation, and
                                               ***herpes***
                                                                     ***simplex***
     infection.
                (3) one or more oxyradical inhibitors, such as desferrioxamine
ΜМ
     or allopurinol, and/or one or more biological modifier agents, such as ***calpain*** inhibitors. The formulations of these compositions is
     dependent upon the specific pathological condition sought to be treated
     or prevented, the.
                active ingredients, typically selected from the group
TD
     consisting of: N-2-mercaptopropionylglycine, N-acetylcysteine,
     glutathione, dimethyl thiourea, desferrioxamine, mannitol,
     plasminogen activator, streptokinase, urokinase, nonsteroidal
     anti-inflammatory agent, cortisone, and carotenoids. Antioxidant
     salen-Mn complexes may also.
   ANSWER 18 OF 26 USPATFULL ON STN
                          1998:58182 USPATFULL
CESSION NUMBER:
                          Lactacystin analogs
TLE:
                          Fenteany, Gabriel, Cambridge, MA, United States
Jamison, Timothy F., Cambridge, MA, United States
VENTOR(S):
                         Schreiber, Stuart L., Boston, MA, United States
Standaert, Robert F., Arlington, MA, United States
President and Fellows of Harvard College, Cambridge,
TENT ASSIGNEE(S):
                          MA, United States (U.S. corporation)
                                NUMBER
                                               KIND
                                                         DATE
                          us 5756764
                                                       19980526
TENT INFORMATION:
                                                       19950606
PLICATION INFO.:
                         us 1995-466468
                                                                   (8)
                          Division of Ser. No. US 1995-421583, filed on 12 Apr
LATED APPLN. INFO.:
                          1995
CUMENT TYPE:
                          Utility
LE SEGMENT:
                          Granted
                          Richter, Johann
IMARY EXAMINER:
SISTANT EXAMINER:
                          Stockton, Laura L.
                          Fish & Richardson P.C.
GAL REPRESENTATIVE:
MBER OF CLAIMS:
                          16
EMPLARY CLAIM:
NE COUNT:
                          2392
S INDEXING IS AVAILABLE FOR THIS PATENT.
     US 5756764
                                  19980526
     The compounds disclosed herein are highly selective for the proteasome,
MM
     and do not inhibit other proteases such as trypsin, .alpha.-
chymotrypsin, ***calpain*** I, ***calpain*** II, papain, and
     cathepsin B.
     . . . the X/MB1 subunit and .alpha.-chain of the proteasome and do not inhibit the activity of proteases such as trypsin, .alpha.-chymotrypsin, ***calpain*** I, ***calpain*** II, cathepsin, and papain. Such selectivity is useful to formulate a
MM
     pharmaceutical composition with fewer side effects and to evaluate.
ММ
     Other eukaryotic transcription factors that require proteolytic
     processing include the general transcription factor TFIIA,

***herpes*** ***simplex*** virus VP16 accessory pr
     TD
     Tris-HCL, pH 8, 20 mM CaCl.sub.2 (plus 100 .mu.M_Cbz-GGR-.beta.NA_for
                 ***Calpain*** I: 20 mM Tris-HCL, pH 8, 1 mM CaCl.sub.2, 1 mM 100 .mu.M Suc-LLVY-AMC for assay); ***Calpain*** II: 20 mM
     DTT (plus 100 .mu.M Suc-LLVY-AMC for assay);
     Tris-HCL, pH 8, 10 mM CaCl.sub.2, 1 mM DTT (plus 100 .mu.M Suc-LLVY-AMC
```

```
ETD
                      TABLE 4
nhibition of Other Proteases
              Effect of lactacystin (100
rotease tested
              .mu.M)
alpha.-Chymotrypsin
              No inhibition
rypsin
              No inhibition
   ***Calpain***
                           No inhibition
                    Τ
   ***Calpain***
                     II
                           No inhibition
              No inhibition
apain
              No inhibition
athepsin B
                containing the human p105 cDNA. Forty-eight hrs after
ETD
      transfection, cells were pretreated for 1 hour with 0.5% DMSO, 50 .mu.M
        ***calpain***
                          inhibitor II, 50 .mu.M MG132, or 10 .mu.M
      beta.-lactone. Cells were then pulse labelled with 250 uCi/plate of.
      .sup.35 S-methionine/cysteine.
      apparent, as was expected (Fan and Maniatis, 1991, Nature 354:395; Palombella et al., 1994, Cell 78:773). Pretreatment of cells with ***calpain*** inhibitor II has no effect on the processing of
ETD
      p105 to p50 (lane 4). However, treatment of cells with the.
    ANSWER 19 OF 26 USPATFULL ON STN
CCESSION NUMBER:
                         1998:57716 USPATFULL
                         Aptamers specific for biomolecules and methods of
ITLE:
                         making
                         Griffin, Linda, Atherton, CA, United States
NVENTOR(S):
                         Albrecht, Glenn, Redwood City, CA, United States
                         Latham, John, Palo Alto, CA, United States
                         Leung, Lawrence, Hillsborough, CA, United States
                         Vermaas, Eric, Oakland, CA, United States
                         Toole, John J., Burlingame, CA, United States
Gilead Sciences, Inc., Foster City, CA, United States
ATENT ASSIGNEE(S):
                          (U.S. corporation)
                               NUMBER
                                              KIND
                                                       DATE
                                                    19980526
                         us 5756291
ATENT INFORMATION:
                                                    19950607
                         us 1995-484192
PPLICATION INFO.:
                         Continuation of Ser. No. US 1992-934387, filed on 21
ELATED APPLN. INFO.:
                         Aug_1992, now abandoned
OCUMENT TYPE:
                         Utility
ILE SEGMENT:
                         Granted
RIMARY EXAMINER:
                         Zitomer, Stephanie W.
                         Bosse, Mark L.
EGAL REPRESENTATIVE:
UMBER OF CLAIMS:
                         12
XEMPLARY CLAIM:
                         6 Drawing Figure(s); 6 Drawing Page(s)
UMBER OF DRAWINGS:
                         8242
INE COUNT:
AS INDEXING IS AVAILABLE FOR THIS PATENT.
      us 5756291
                                 19980526
                                                                            <--
                                (HA)
ETD
euraminidase (NA)
ucleoprotein (NP)
1 and M2 proteins
S1 and NS2 proteins
epatitis B
nvelope (surface antigenP proteins (including pre-S1,
re-S2 and S)
ucleocapsid (core) proteins
-gene product
-gene product
    ***Cytomegalovirus***
mmediate early (alpha) gene products (including IE1 and
E2)
arly (beta) gene products (including DNA pol pl40, DBP52 DBP 140)
```

irus-encoded envelope glycoproteins

hymidine kinase

pstein-Barr Virus

ibonucleotide reductase

```
RLF1 protein)
early gene products (including SMLF1, MRF1, ALF2,. . . synthase
alanine aminotransferase
alcohol dehydrogenase
aldolase
adose reducase
alkaline phosphatase
amidophosphodbosylanine transferase
AMP phosphodiesterase
amyloid b/A4 protein
amyloid precursor protein
ankarin
arginase
argininosuccinate lyase
argininosuccinate synthetase
aromatase
aryl sulfatase
aspartate aminotransferase
aspartate transcarbamoylase
ATP diphosphohydrolase
ATPase
b-actin
b-glucuronidase
b-glycerophosphatase
b-ketoacyl-ACP reductase
b-ketoacyl-ACP sythetase
b-spectrin
b-tropomyosin
b-tubulin
C5a inactivation factor
calcitoin
calmodulin
    ***calpain***
calreticulin
carbamoyl-phosphate synthetase
carbonic anhydrase
casein kinase 1
casein kinase 2
catalase
catechol methyltransferase
cathepsin
cathepsin B and L
cdc 2 p34
cdc 10
cdc 13 p60
cdc 25 p80
chaparonin
cholesterol esterase
cholesterol monooxygenase
citrate.
     ANSWER 20 OF 26 USPATFULL on STN
1.6
ACCESSION NUMBER:
                          1998:24868
                                      USPATFULL
TITLE:
                          Non-crosslinked protein particles for therapeutic and
                          diagnostic use
INVENTOR(S):
                          Yen, Richard C. K., Yorba Linda, CA, United States
PATENT ASSIGNEE(S):
                          Hemosphere, Inc., Irvine, CA, United States (U.S.
                          corporation)
                                NUMBER
                                              KIND
                                                       DATE
                          us 5725804
                                                     19980310
PATENT INFORMATION:
APPLICATION INFO.:
                          US 1995-471650
                                                     19950606
                                                                (8)
RELATED APPLN. INFO.:
                          Continuation-in-part of Ser. No. US 1994-212546, filed
                          on 14 Mar 1994, now patented, Pat. No. US 5616311 which
                          is a continuation-in-part of Ser. No. US 1993-69831,
                          filed on 1 Jun 1993, now abandoned And Ser. No. US
                          1992-959560, filed on 13 Oct 1992, now patented, Pat. No. US 5308620 which is a continuation-in-part of Ser.
                          No. US 1991-641720, filed on 15 Jan 1991, now abandoned
                          Utility
DOCUMENT TYPE:
FILE SEGMENT:
                          Granted
                          Lovering, Richard D.
Townsend & Townsend & Crew
PRIMARY EXAMINER:
LEGAL REPRESENTATIVE:
NUMBER OF CLAIMS:
                          11
EXEMPLARY CLAIM:
                          1
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CAS INDEXING IS AVAILABLE FOR THIS PATENT.
        US 5725804
***Calpain*** I
PΙ
                                      19980310
                               Inhibitor I
Inhibitor II
DETD
           ***Calpain***
DETD
           ***Calpain*** Inhibitor Peptide
DETD
DETD
                   double stranded), cloning vectors, coliphage DNA, lambda phage
        DNA, M13 DNA, Adenovirus DNA, phi-X 174 phage DNA, Simian virus DNA,
           ***Cytomegalovirus*** DNA, Epstein-Barr Virus genes, ***Herpes***
                              genes, ribosomal RNA, human DNA and RNA; Genes coding
           ***Simplex***
        for ribozymes; genes coding for antibiotics (e.g., ampicillin,
         chloramphenicol, cycloserine, gentamycin, . .
     ANSWER 21 OF 26 USPATFULL ON STN
                             97:115268 USPATFULL
ACCESSION NUMBER:
                              Synthetic catalytic free radical scavengers useful as
TITLE:
                              antioxidants for prevention and therapy of disease
INVENTOR(S):
                              Malfroy-Camine, Bernard, Arlington, MA, United States
                              Doctrow, Susan Robin, Roslindale, MA, United States
PATENT ASSIGNEE(S):
                             Eukarion, Inc., Bedford, MA, United States (U.S.
                              corporation)
                                   NUMBER KIND
                                                           DATE
                             us 5696109
us 1995-485489
PATENT INFORMATION:
                                                            19971209
                                                            19950607 (8)
APPLICATION INFO.:
                             Continuation-in-part of Ser. No. US 1995-380731, filed
RELATED APPLN. INFO.:
                             on 26 Jan 1995 which is a continuation-in-part of Ser. No. US 1992-987474, filed on 7 Dec 1992, now patented, Pat. No. US 5403834
                                      NUMBER
                                                      DATE
PRIORITY INFORMATION:
                             wo 1993-us11857 19931206
DOCUMENT TYPE:
                             Utility
FILE SEGMENT:
                             Granted
                             Jarvis, William R. A.
PRIMARY EXAMINER:
LEGAL REPRESENTATIVE:
                              Townsend and Townsend and Crew LLP
NUMBER OF CLAIMS:
                              14
EXEMPLARY CLAIM:
NUMBER OF DRAWINGS:
                             28 Drawing Figure(s); 19 Drawing Page(s)
LINE COUNT:
                             3441
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
ΡI
        us 5696109
                                      19971209
        . . . effects in treating systemic lupus erythematosus, Crohn's disease, gastric ulcers, oxygen toxicity, burned patients, renal failure attendant to transplantation, and ***herpes*** ***simplex***
SUMM
        infection.
SUMM
                   (3) one or more oxyradical inhibitors, such as desferrioxamine
        or allopurinol, and/or one or more biological modifier agents, such as ***calpain*** inhibitors. The formulations of these compositions is
                              inhibitors. The formulations of these compositions is
        dependent upon the specific pathological condition sought to be treated
        or prevented, the. . . . . active ingredients, typically selected from the group consisting of: N-2-mercaptopropionylglycine, N-acetylcysteine, glutathione, dimethyl thiourea, desferrioxamine, mannitol, .alpha.-tocopherol, ascorbate, allopurinol, 21-aminosteroids, ***calpain*** inhibitors, glutamate receptor antagonists, tplasminogen activator, streptokinase, urokinase, nonsteroidal
DETD
        anti-inflammatory agent, cortisone, and carotenoids. Antioxidant
        salen-Mn complexes may also.
     ANSWER 22 OF 26 USPATFULL on STN
L6
                             97:26904 USPATFULL
ACCESSION NUMBER:
TITLE:
                             Non-crosslinked protein particles for therapeutic and
                             diagnostic use
INVENTOR(S):
                             Yen, Richard C. K., Glendora, CA, United States
PATENT ASSIGNEE(S):
                             Hemosphere, Inc., Irvine, CA, United States (U.S.
                             corporation)
                                   NUMBER KIND DATE
                             us 5616311
PATENT INFORMATION:
                                                           19970401
APPLICATION INFO.:
                             us 1994-212546
                                                           19940314 (8)
                             Continuation-in-part of Ser. No. US 1993-69831, filed
RELATED APPLN. INFO.:
                             on 1 Jun 1993, now abandoned And Ser. No. US 1992-959560, filed on 13 Oct 1992, now patented, Pat.
```

```
No. US 1991-641720, filed on 15 Jan 1991, now abandoned
                            Utility
DOCUMENT TYPE:
FILE SEGMENT:
                            Granted
                            Lovering, Richard D.
Townsend & Townsend & Crew
PRIMARY EXAMINER:
LEGAL REPRESENTATIVE:
NUMBER OF CLAIMS:
                            26
EXEMPLARY CLAIM:
                            1,26
LINE COUNT:
                            2585
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
       US 5616311
                                    19970401
ΡI
                                                                                 <--
                             Inhibitor I
          ***Calpain***
DETD
          ***Calpain***
                             Inhibitor II
Inhibitor Peptide
DETD
          ***Calpain***
DETD
                  double stranded), cloning vectors, coliphage DNA, lambda phage
DETD
       DNA, M13 DNA, Adenovirus DNA, phi-X 174 phage DNA, Simian virus DNA, ***Cytomegalovirus*** DNA, Epstein-Bart Virus genes, ***Herpes
          ***Simplex***
                            genes, ribosomal RNA, human DNA and RNA; Genes coding
        for ribozymes; genes coding for antibiotics (e.g., ampicillin,
        chloramphenicol, cycloserine, gentamycin,.
     ANSWER 23 OF 26 USPATFULL on STN
L6
                            95:29636 USPATFULL
ACCESSION NUMBER:
TITLE:
                            Synthetic catalytic free radical scavengers useful as
                            antioxidants for prevention and therapy of disease
INVENTOR(S):
                            Malfroy-Camine, Bernard, Arlington, MA, United States
                            Baudry, Michel, Long Beach, CA, United States
Eukarion, Inc., Arlington, MA, United States (U.S.
PATENT ASSIGNEE(S):
                            corporation)
                                 NUMBER
                                                KIND
                                                          DATE
                            -----
                            us 5403834
PATENT INFORMATION:
                                                         19950404
                                                                                 <--
                            US 1992-987474
APPLICATION INFO.:
                                                        19921207 (7)
DOCUMENT TYPE:
                            Utility
FILE SEGMENT:
                            Granted
                            Henley, III, Raymond Criares, T. J.
PRIMARY EXAMINER:
ASSISTANT EXAMINER:
LEGAL REPRESENTATIVE:
                            Dunn, Tracy J.
NUMBER OF CLAIMS:
EXEMPLARY CLAIM:
NUMBER OF DRAWINGS:
                            9 Drawing Figure(s): 6 Drawing Page(s)
LINE COUNT:
                            1742
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
PΙ
       US 5403834
                                    19950404
       . . . effects in treating systemic lupus erythematosus, Crohn's disease, gastric ulcers, oxygen toxicity, burned patients, renal failure attendant to transplantation, and ***herpes*** ***simplex***
SUMM
       infection.
SUMM
                  (3) one or more oxyradical inhibitors, such as desferrioxamine
       or allopurinol, and/or one or more biological modifier agents, such as ***calpain*** inhibitors. The formulations of these compositions is
       dependent upon the specific pathological condition sought to be treated
       or prevented, the. . . . . active ingredients, typically selected from the group consisting of: N-2-mercaptopropionylglycine, N-acetylcysteine,
DETD
       glutathione, dimethyl thiourea, desferrioxamine, mannitol
        .alpha.-tocopherol, ascorbate, allopurinol, 21-aminosteroids,
       ***calpain*** inhibitors, glutamate receptor antagonists, tissue plasminogen activator, streptokinase, urokinase, nonsteroidal
          ***calpain***
       anti-inflammatory agent, cortisone, and carotenoids. Antioxidant
       salen-Mn complexes may also.
     ANSWER 24 OF 26 USPATFULL on STN
L6
ACCESSION NUMBER:
                            94:51514 USPATFULL
TITLE:
                            Antiplatelet and antithrombotic activity of platelet
                            glycoprotein Ib receptor fragments
INVENTOR(S):
                            Handin, Robert, Needham, MA, United States
                            Brigham and Women's Hospital, Boston, MA, United States
PATENT ASSIGNEE(S):
                            (U.S. corporation)
                                 NUMBER
                                               KIND
                                                        DATE
PATENT INFORMATION:
                           US 5321127
                                                        19940614
                                                                                 <--
APPLICATION INFO.:
                           us 1991-670606
                                                        19910318 (7)
DOCUMENT TYPE:
                           Utility
FILE SEGMENT:
                           Granted
```

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MBER OF CLAIMS:
EMPLARY CLAIM:
MBER OF DRAWINGS:
                         10 Drawing Figure(s); 8 Drawing Page(s)
                         1494
NE COUNT:
S INDEXING IS AVAILABLE FOR THIS PATENT.
                                 19940614
               et al., Blood 67:19-26 (1986)). It is cleaved from the surface
MM
        intact platelets by various maneuvers which activate platelet
       ***calpain***
                         , an endogenous calcium dependent protease (Fox,
     et al., J. Biol. Chem. 263:4882-4890 (1988)). Further digestion of
     glycocalicin with.
                at the Massachusetts General Hospital, Boston Mass. to produce
TD
     pCDM8-GpIba. The CDM8 vector contains a cloning site downstream from the ***cytomegalovirus*** promoter as well as the SV40 origin of
     replication, permitting transient expression of the heterologous protein
     in COS cells (Aruffo,.
   ANSWER 25 OF 26 BIOSIS COPYRIGHT 2004 BIOLOGICAL ABSTRACTS INC. on STN
                    1999:184941 BIOSIS
CESSION NUMBER:
CUMENT NUMBER:
                    PREV199900184941
                             ***cytomegalovirus*** -activated
                                                                       ***calpain***
TLE:
                    Human
                    and p21Cip1 degradation in human lung fibroblasts.
THOR(S):
                    Chen, Z.; Knutson, E.; Kurosky, A.; Liu, S.; Albrecht, T.
RPORATE SOURCE:
                    Univ. Texas Med. Branch, Galveston, TX 77555, USA
                    Proceedings of the American Association for Cancer Research
Annual Meeting, (March, 1999) Vol. 40, pp. 447-448. print.
Meeting Info.: 90th Annual Meeting of the American
URCE:
                    Association for Cancer Research. Philadelphia,
Pennsylvania, USA. April 10-14, 1999. American Association
                    for Cancer Research.
                    ISSN: 0197-016x
                    Conference; (Meeting)
CUMENT TYPE:
                    Conference; Abstract; (Meeting Abstract)
                    English
NGUAGE:
                    Entered STN: 5 May 1999
Last Updated on STN: 5 May 1999
TRY DATE:
            ***cytomegalovirus*** -activated
                                                     ***calpain***
                                                                        and p21Cip1
   degradation in human lung fibroblasts.
   Proceedings of the American Association for Cancer Research Annual
   Meeting, (March, 1999) Vol. 40, pp. 447-448. print.
   Meeting Info.: 90th Annual Meeting of the American Association for Cancer
   Research. Philadelphia, Pennsylvania, USA. April 10-14, 1999. American
   Association for Cancer Research.
   ISSN: 0197-016x.
   Major Concepts
      Tumor Biology
   Parts, Structures, & Systems of Organisms
      lung fibroblasts
  Chemicals & Biochemicals
___***calpain*** ; p21-Cip1: degradation
GN Classifier
      Herpesviridae
                        03115
   Super Taxa
      dsDNA Viruses; Viruses; Microorganisms
               ***cytomegalovirus*** : pathogen
      human
  Taxa Notes
      Double-Stranded DNA Viruses, Microorganisms, Viruses
GN Classifier
      Hominidae
                    86215
   Super Taxa
      Primates; Mammalia; Vertebrata; Chordata; Animalia
   Organism.
                  ***calpain*** )
   78990-62-2 (
  ANSWER 26 OF 26 BIOSIS COPYRIGHT 2004 BIOLOGICAL ABSTRACTS INC. on STN SSION NUMBER: 1993:407472 BIOSIS MENT NUMBER: PREV199396073197
CESSION NUMBER:
CUMENT NUMBER:
TLE:
                    Inhibition of proteolytic activity of poliovirus and
                    rhinovirus 2A proteinases by elastase-specific inhibitors.
THOR(S):
                    Molla, Akhteruzzaman; Hellen, Christopher U. T.; Wimmer,
                    Eckard [Reprint author]
                    Dep. Microbiol., Sch. Med., State Univ. New York at Stony Brook, Stony Brook, NY 11794-8621, USA
RPORATE SOURCE:
JRCE:
                    Journal of Virology, (1993) Vol. 67, No. 8, pp. 4688-4695.
```

Sterne, Kessler, Goldstein & Fox

GAL REPRESENTATIVE:

```
MENT TYPE:
                      Article
                      English
UAGE:
 DATE: Entered STN: 8 Sep 1993
Last Updated on STN: 6 Nov 1993
Journal of Virology, (1993) Vol. 67, No. 8, pp. 4688-4695.
CODEN: JOVIAM. ISSN: 0022-538X.
Y DATE:
         was unchanged, an observation suggesting that this inhibitor may
 have formed a covalent bond with the active-site Cys-109 nucleophile. Iodoacetamide, ***calpain*** inhibitor 1, and antipain inhibited poliovirus 2A-pro. MPCMK caused a reduction in the yields of the
 enteroviruses poliovirus type 1.
 Major Concepts
     Enzymology (Biochemistry and Molecular Biophysics); Microbiology;
     Pharmacology
 Chemicals & Biochemicals
     PROTEINASES; ELASTASE; ELASTATINAL; IODOACETAMIDE;
                                                                                 ***CALPAIN***
     ANTIPAIN
 Miscellaneous Descriptors
     CORNEAL INFLAMMATION; GRANULOCYTE-MACROPHAGE COLONY STIMULATING FACTOR; ***HERPES*** ***SIMPLEX*** VIRUS-TYPE I; INTERLEUKIN-1-ALPHA;
     INTERLEUKIN-10; INTERLEUKIN-4; INTERLEUKIN-6
 9001-92-7D (PROTEINASES)
 9004-06-2 (ELASTASE)
 51798-45-9 (ELASTATINAL)
 144-48-9 (IODOACETAMIDE)
 78990-62-2 ( ***CALPAIN***
37691-11-5 (ANTIPAIN)
ogging off of STN---
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SINCE FILE

SINCE FILE

ENTRY

**ENTRY** 

-2.77

176.82

TOTAL SESSION

177.24

TOTAL SESSION

-2.77

cuting the logoff script...

COUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)

INTERNATIONAL LOGOFF AT 13:39:02 ON 08 JUN 2004

IN U.S. DOLLARS

. ESTIMATED COST

SUBSCRIBER PRICE

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